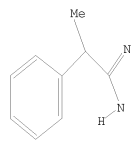
7 8 9 12 ring nodes : 1 2 3 4 5 6 ring/chain nodes : 10 11 chain bonds : 5-7 7-8 7-9 9-10 9-11 11-12 ring bonds : 1-2 1-6 2-3 3-4 4-5 5-6 exact/norm bonds : 9-10 9-11 exact bonds : 5-7 7-8 7-9 11-12 normalized bonds : 1-2 1-6 2-3 3-4 4-5 5-6 isolated ring systems : containing 1 :

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS 11:CLASS

L1 STRUCTURE UPLOADED

=> d 11 L1 HAS NO ANSWERS L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> s 11 SAMPLE SEARCH INITIATED 10:41:11 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 360 TO ITERATE

100.0% PROCESSED 360 ITERATIONS 9 ANSWERS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 6062 TO 8338

Habte 06/05/2008

PROJECTED ANSWERS: 9 TO 360

L2 9 SEA SSS SAM L1

=> s l1 sss full

FULL SEARCH INITIATED 10:41:36 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 6833 TO ITERATE

100.0% PROCESSED 6833 ITERATIONS 137 ANSWERS

SEARCH TIME: 00.00.01

L3 137 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST 178.36 178.57

FILE 'CAPLUS' ENTERED AT 10:41:42 ON 05 JUN 2008
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 5 Jun 2008 VOL 148 ISS 23 FILE LAST UPDATED: 4 Jun 2008 (20080604/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

http://www.cas.org/legal/infopolicy.html

=> s 13

L4 60 L3

=> d ibib abs hitstr tot

Habte 06/05/2008

Page 5 10/568,760

L4 ANSWER 1 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2007:1303013 CAPLUS DOCUMENT NUMBER: 147:541746
Preparation of quinolinones and analogs as antiviral agents
Kumar, Dange V.; Rai, Roopa; Young, Wendy B.; Hu,
Huiyong; Riggs, Jennifer R.; Ton, Tony Loc; Green,
Michael J.; Hart, Barry P.; Brameld, Kenneth A.;
Dener, Jeff M.
Virobay, Inc., USA
PCT Int. Appl., 201pp.
CODEN: PIXXD2 147:541746 TITLE: INVENTOR(S): PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT	NO.		KINI	_	DATE				ICAT					ATE	
WO 2007			A2		2007				007-					0070	
WO 2007	130499		A3		2008	0110									
W:	AE, AG	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,	CA,
	CH, CN	. co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,
	GD, GE	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,
	KN, KP	KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	ME,	MG,
	MK, MN	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,
	RO, RS	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	TJ,	TM,	TN,	TR,
	TT, TZ	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	zw					
RW:	AT, BE	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
	IS, IT	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,
	BJ, CF	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,
	GH, GM	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,
	BY, KG	KZ,	MD,	RU,	TJ,	TM,	AP,	EA,	EP,	OA					
US 2007	US 20070287699					1213		US 2	007-	7424	51		21	0070	430
PRIORITY APP	LN. INF).:						US 2	006-	7969	43P	1	P 21	0060	501

OTHER SOURCE(S): MARPAT 147:541746

ANSWER 1 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) Benzeneethanimidamide, 4-fluoro-N-hydroxy- α , α -dimethyl- (CA

L4 ANSWER 1 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

Title compds. represented by the formula I & II & III [wherein Zl = N or CR1; Z2 = N or CR2; Z3 = N or CR4; R1 = H, halo, alkyl, etc.; R2 = H, halo, alkoxy, etc.; R3 = halo, alkyl, aryl, etc.; R4 = H, halo,

haloalkyl, etc., ...
haloalkyl, etc., R5 = alkyl, cycloalkylamino, arylamino, etc.; R6 = (un)substituted
1,2,4-oxadiazol-3-yl, 1,2,4-oxadiazol-5-yl, 1,3,4-oxadiazol-2-yl or
oxazol-2-yl; and pharmaceutically acceptable salts thereof) were

oxazol-z-y1; and pharmaceutically acceptable saits thereof) were ared as antiviral agents. For example, IV was provided in a multi-step synthesis starting from reaction of Et 2-cyano-3-ethoxyacrylate with 3,4-dimethoxyphenylamine. The invention compds. showed activity in HCV replicon assays and their formulations were also presented. Thus, the title compds. and their pharmaceutical compns. are useful for the treatment of viral infections, particularly HCV.
957140-71-5, N-Hydroxy-2-methyl-2-phenylpropanimidamide
8L: NCT (Reactant); RACT (Reactant or reagent)
(preparation of 4-quinolinones and analogs as antiviral agents)
957140-71-5 CAPLUS
Benzeneethanimidamide, N-hydroxy-α,α-dimethyl- (CA INDEX NAME)

RN 957140-73-7 CAPLUS

ANSWER 2 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 2007:1050755 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR(S):

2007:1059755 CAPLES
2007:1059755 CAPLES
2007:1059755 CAPLES
2007:1059755 CAPLES
2007:1059755 CAPLES
2007:1059755 CAPLES
2007:105975 CAPLES
2007:10

CORPORATE SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

AB The preparation and structure-activity relationships of novel pyrrolidine-carboxamides and oxadiazoles are described. Compds. in this series were found to be potent hNKI antagonists in vitro and efficacious in vivo with minimal interactions with P450 liver enzymes. Oxadiazole analog (I) was determined to have excellent hNKI binding affinity, functional activity, and a good PD response in vivo.

IT 957476-30-IP RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (pyrrolidine-carboxamides and oxadiazoles as potent hNKI antagonists)

RN 957476-30-I CAPIUS

CN Benzeneethanimidamide, N-hydroxy-α,α-dimethyl-3,5-bis(trifluoromethyl)- (CA INDEX NAME)

ANSWER 2 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

REFERENCE COUNT: THERE ARE 12 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ACCESSION NUMBER:

DOCUMENT NUMBER:

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
GI

TITLE: AUTHOR(S): CORPORATE SOURCE:

SOURCE:

Novel triazine analogs of 5-alkyl-2-alkylthio-6-[1-(2,6-difluorophenyl)alkyl]-3,4-dihydropyrimidin-4(3H)-ones (F2-DABOs), previously described by us as nonnucleoside HIV-1 reverse transcriptase inhibitors (NNRTIS), were tested for their antiproliferative and cytodifferentiating activity on the A-375 human melanoma cell line. Most of the tested derivs. were effective in decreasing cell proliferation, facilitating morphol. differentiation, and reprogramming gene expression. All these effects were reversible upon withdrawal of RT inhibitors.

ANSWER 3 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

146:454167

2007:344613 CAPLUS

6-alkylthio-4-[1-(2,6-difluorophenyl)alkyl]-1H-

The compds. tested, 3f $\left(\mathrm{I}\right)$ showed the highest antiproliferative effect, whereas compound 6c, although not affecting cell proliferation, is

wed with a strong cytodifferentiating effect, which is probably related to a marked upregulation of the e-cad gene. These results support the potential of NNRTIs as valuable antitumor agents.

935480-63-UP

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(triazinones as regulators of cell differentiation and proliferation)

ANSWER 3 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN 935480-63-0 CAPLUS (Continued)

nzeneethanimidamide, 2,6-difluoro-α-methyl- (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 47 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

INVENTOR(S):

2007:174406 CAPLUS
146:251662
146:251662
Xanthine derivatives as selective HM74A agonists and their preparation
Hatley, Richard Jonathan Daniel; Heer, Jag Paul;
Liddle, John, Mason, Andrew Memurtrie; Pinto, Ivan
Leo; Rahman, Shahzad Sharooq; Smith, Ian Edward David
Smithkline Beecham Corporation, USA
PCT Int. Appl., 312pp.
CODEN: PIXXD2
Patent
English
2

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE EP 1912992 A1 20080423 EP 2000-776059 200800000

R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FF, GB, GR, HU, IE,

IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, HR

KR 2008038396 A 20080506 KR 2008-705724 20080307

KITY APPLN. INFO:: GB 2005-16464 A 20550810 PRIORITY APPLN. INFO.: GB 2006-7736 A 20060419 GB 2006-14569 A 20060721 WO 2006-EP7869 W 20060808

OTHER SOURCE(S): MARPAT 146:251662

06/05/2008 Habt.e

10/568,760

Page 7

L4 ANSWER 4 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

AB The invention relates to compds. of formula I, which are xanthine derivs.,

The invention relates to compds. of formula I, which are xanthine derivs., processes for the manufacture of said derivs., pharmaceutical formulations containing the active compds. and the use of the compds. in therapy, for example, in the treatment of diseases where under-activation of the BM74A receptor contributes to the disease or where activation of the receptor will be beneficial. Compds. of formula I wherein Rl is (un)substituted C1-10 alkylene; R2 is B, (un)substituted C1-10 alkyln, (un)substituted C2-10

alkenyl, (un)substituted C2-10 alkynyl, (un)substituted cycloalkyl, (un)substituted cycloalkenyl, (un)substituted heterocyclyl, and (un)substituted (hetero)aryl; R3 is halo and CN; and their pharmaceutically acceptable derivs. thereof, are claimed. Example

compound
If was prepared by alkylation of 8-chloro-3-pentyl-7-(2-propen-1-y1)-3,7dihydro-1H-purine-2,6-dione with 2-chloroethanol followed by

deallylation.
All the invention compds. were evaluated for their HM74A agonistic

activity.

925444-91-3P 925444-98-0P
RL: IMP (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RRCT (Reactant or reagent) (Intermediate; preparation of xanthine derive. as selective HM74A

agonists)
RN 925444-91-3 CAPLUS
CN Benzenethanimidamide, 3-bromo-N-hydroxy-α,α-dimethyl-,
[C(Z)]- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \quad \text{Me} \\ \text{Br} \\ \text{Z} \\ \text{OH} \\ \text{NH}_2 \end{array}$$

925444-98-0 CAPLUS Benzeneethanimidamide, N,4-dihydroxy- α , α -dimethyl-, [C(Z)]-

ANSWER 4 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

L4 ANSWER 4 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) (CA INDEX NAME)

Double bond geometry as shown.

42191-51-5 925698-75-5 925893-03-4
RL: RCT (Reactant); RACT (Reactant or reagent)
(starting material; preparation of xanthine derivs. as selective HM74A agonists)
42191-51-5 CAPLUS
Benzeneethanimidamide, N-hydroxy-α-methyl- (CA INDEX NAME)

925698-75-5 CAPLUS Benzeneethanimidamide, N-hydroxy- α -methyl-, [C(Z)]- (CA INDEX NAME)

Double bond geometry as shown.

925893-03-4 CAPLUS Benzeneethanimidamide, 4-chloro-N-hydroxy- α -methyl-, hydrochloride (1:1) (CA INDEX NAME)

Double bond geometry as shown.

L4 ANSWER 5 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 2007:174405 CAPLUS
DOCUMENT NUMBER: 146:251661
TITLE: Preparation of xanthine derivatives as selective

agonists

Batley, Richard Jonathan Daniel; Mason, Andrew Mcmurtrie; Pinto, Ivan Leo
Smithkline Beecham Corporation, USA
PCT Int. Appl., 199pp.
CODEN: PIXXD2
Patent
English
2 INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA	PATENT NO.										ICAT						
WO	2007									WO 2	006-	EP78	65		2	0060	808
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KΡ,
		KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,
		MW,	MX,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RS,	RU,
		SC,	SD,	SE,	SG,	SK,	SL,	SM,	SY,	ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,
		US,	UZ,	VC,	VN,	ZA,	ZM,	zw									
	RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
		IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,
		CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,
		GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
		KG,	KZ,	MD,	RU,	TJ,	TM										
AU	2006	2782	15		A1		2007	0215		AU 2	006-	2782	15		2	0060	808
EP	1912	991			A1		2008	0423		EP 2	006-	7630	16		2	0060	808
	R:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
											PT,						
	2008				A												
PRIORIT	Y APP	LN.	INFO	. :						GB 2	005-	1646	4		A 2	0050	810
										GB 2	006-	7736			A 2	0060	419
										GB 2	006-	1456	9		A 2	0060	721
										WO 2	006-	EP 78	65		W 2	0060	808

OTHER SOURCE(S): MARPAT 146:251661

06/05/2008 Habte

ANSWER 5 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) Xanthine derivs. of formula I [R1 = (CH2)mX(CH2)nY; X = heteroaryl, heterocyply]; Y = (substituted) aryl, heteroaryl, aryloxy; m = 3-4; n = 0-1; R2 = (substituted) alkyl; R3 = halo] are prepared for the treatment

diseases where under-activation of the HM74A receptor contributes to the disease or where activation of the receptor will be beneficial. Thus, I

disease or where activation of the receptor will be beneficial. Thus, II was prepared from ntyl-8-chloro-7-allyl-3,7-dihydro-1H-purine-2,6-dione, 4-(3-hydroxypropyl)pyrazole and 2-chloro-6-fluorobenzyl bromide. The prepared compds. had pEC50 values ≥ 4.3 and efficacy ≥ 30% in GTPyS binding assays. 925444-91-3 925444-98-0P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of xanthine derivs. as selective HM74A agonists) 925444-91-3 CAPLUS Benzeneethanimidamide, 3-bromo-N-hydroxy-α,α-dimethyl-, [C(Z)]- (CA INDEX NAME)

Double bond geometry as shown.

925444-98-0 CAPLUS Benzeneethanimidamide, N,4-dihydroxy- α , α -dimethyl-, [C(Z)]-(CA INDEX NAME)

Double bond geometry as shown.

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

REFERENCE COUNT:

ANSWER 6 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)
RENCE COUNT: 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 6 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:30368 CAPLUS 146:317014

DOCUMENT NUMBER:

146:317014

New Optically Active N-Heterocyclic Carbene Complexes for Hydrogenation: A Tale with an Atropisomeric Twist Chen, Dianjun; Bamphavichit, Vorawit; Reibenspies, Joe; Burgess, Kevin Department of Chenistry, Texas A and M University, College Station, TX, 77843, USA
Organometallics (2007), 26(4), 855-859
CODEN: ORGMD7; ISSN: 0276-7333
American Chemical Society
Journal TITLE: AUTHOR(S):

CORPORATE SOURCE:

SOURCE

DIEBLISHER.

PUBLISHER: American Chemical Society
DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 146:317014
AB Iridium chiral 1,2,4-triazol-3-ylidene pyrimidine complexes derived from isoleucine were prepared and examined as catalysts for asym. hydrogenation of
1,2-diphenyl-1-propene; the complexes low enantioselectivity of 12% ee. Homologation and alkynylation of L-isoleucine afforded
(33,4R)-4-Boc-amino-3-methyl-7-dodecyn-6-one (8), which was condensed with

anddines R(NHN)M12 to give N-Boc-protected (α N) -2-K-6-Bu- α - [(13)-1-methylpropyl]-4-pyrimidineethanamines (9a-c; R = Ph, 1-adamantyl, CMePh2). Deprotection of 9a-c followed by reaction with 3-(1-adamantyl)-1,3-4-oxadiazolium tetrafluoroborate gave the ligand precursors, 1-adamantyl-4-[(1R,2S)-1-(2-R-6-butyl-4-pyrimidinylmethyl)-2-methylbutylletrazolium tetrafluoroborates (2a-c; R = Ph, 1-adamantyl, CMePh2), which upon metalation and halogen abstraction afforded the corresponding cationic iridium carbene-pyrimidine chelate cyclooctaddene complexes (5). A structure of the 1,2,4-triazolium salts is easily varied, allowing an access to a diverse set of N-heterocyclic carbene complexes. A coordinated chlorine atom was retained on reaction of 2

with [Ir(COD)Cl]2, and this resulted in two atropoisomeric complexes, 3 and 4, which were both characterized via x-ray diffraction studies. Neither of which were both characterized in two atropologments complexes, 3 and 4, which were both characterized via x-ray diffraction studies. Neither of these complexes mediated hydrogenation of (E)-1,2-diphenyl-1-propene, but both 3 and 4 were reacted with NaBAFF4 to give the chlorine-free complex 5, which was catalytically active in this reaction. 173601-37-1

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of chiral carbene iridium triazolylidene-pyrimidine

complexes as catalysts for asym. hydrogenation of alkenes) 173601-37-1 CAPLUS Benzeneethanimidamide, α -methyl- α -phenyl- (CA INDEX NAME)

ANSWER 7 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

2006:189333 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

L4 ANSWER 7 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 2006:189333 CAPLUS
DOCUMENT NUMBER: 146:228672
TITLE: 1-halcalk-1-enes
AUTHOR(S): Schantl, J. G.
CORPORATE SOURCE: Germany
CODEN: SCUURCE: Science of Synthesis (2006), Volume Date 2005, 24, 223-224
CODEN: SSCUY9
PUBLISHER: Georg Thieme Verlag
DOCUMENT TYPE: Journal; General Review
LANGUAGE: English
AB A review of methods to prepare 1-nitrogen-functionalized
1-halcalk-1-enes
1 40645-76-9 English
RL: RCT (Heactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Feactant or reagent)
(review preparation of nitrogen functionalized halcalkenes)
RN 40645-76-9 CAPLUS
CN Benzeneethanimidamide, N-(1-chloro-2-phenyl-1-propen-1-y1)-α-methyl-, hydrochloride (1:1) (CA INDEX NAME)

● HCl

THERE ARE 154 CITED REFERENCES AVAILABLE FOR REFERENCE COUNT: 154 THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT

L4 ANSWER 8 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2005:951698 CAPLUS

DOCUMENT NUMBER:

TITLE:

AUTHOR (S)

CORPORATE SOURCE: SOURCE:

144:467615
Amidines (imidamides) N-substituted by metals, halogens, oxygen, and other heteroatoms Ostrowska, K.; Kolasa, A. Germany Science of Synthesis (2005), 22, 489-563 CODEN: SSCYJ9
Georg Thieme Verlag Journal; General Review English DIIBLISHER DOCUMENT TYPE: LANGUAGE:

UAGE: English
A review of the preparation and synthetic applications of amidine derivs.
160154-90-5P
RL: SFN (Synthetic preparation); PREP (Preparation)
(preparation and synthetic applications of amidine derivs.)
160154-90-5 CAPLUS
Benzeneethanimidic acid, α-hydroxy-α-methyl-, hydrazide (CA
INDEX NAME)

REFERENCE COUNT:

THERE ARE 838 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 9 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

14 ANSWER 9 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER:

DOCUMENT NUMBER:

TITLE:

131.326018

Synthesis and biological activity of flurbiprofen analogues as selective inhibitors of β-amyloidi-42 secretion

AUTHOR(S):

Peretto, Ilaria; Radaelli, Stefano; Parini, Carlo; Zandi, Michele; Raveglia, Luca F.; Dondio, Giulio; Fontanella, Laura; Misiano, Paola; Biogono, Chiara; Rizzi, Andrea; Riccardi, Benedetta; Biscaioli, Marcello; Marchetti, Silvia; Puccini, Paola; Catinella, Silvia; Rondelli, Ivano; Cenacchi, Valentina; Bolzoni, Pier Tonino; Caruso, Paola; Villetti, Gino; Facchinetti, Fabrizio; Del Giudice, Elda; Moretto, Nadia; Imbimbo, Bruno P.

CORPORATE SOURCE:

CORPORATE SOURCE:

ABERIA SALVANIA SA

RE: RET (Reactant); SFN (Synthetic preparation); PREP (Preparation); I (Reactant or reagent) (Synthesis and biol. activity of flurbiprofen analogs as selective inhibitors of \$\beta\$-amploid1-42 secretion devoid of anti-cyclooxygenase activity) 884905-31-1 CAPLUS

[1,1'-Biphenyl]-4-ethanimidamide, 2-fluoro-N-hydroxy-\alpha-methyl-4'-(trifluoromethyl)- (CA INDEX NAME)

L4 ANSWER 9 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 39 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

ANSWER 10 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN 2005:283457

ACCESSION NUMBER: DOCUMENT NUMBER:

2005:283457 CAPLUS
142:355052
Preparation of amidines and their salts useful in the inhibition of chemotaxis of neutrophils induced by interleukin-8
Allegretti, Marcello; Cesta, Maria Candida; Nano, Giuseppe; Bertini, Riccardo; Bizzarri, Cinzia; Colotta, Francesco Dompe S.P.A., Italy
PCT Int. Appl., 22 pp.
CODEN: PIXXD2
Patent
English
1

INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. AU 2004-274183 CA 2004-2539842 CA 2539842 EP 1663960 A1 20050331 20040916 A2 20060607 EP 2004-787150 20040916 EF 1664960 A2 20060601 B EP 2004-787150 C 20040916 EP 2004-787150 EP 200 NO 2006001721 PRIORITY APPLN. INFO.: 20060419 NO 2006-1721 EP 2003-103557

> WO 2004-EP52201 W 20040916

A 20030925

WO 2004-EP52201 W 20040916

HER SOURCE(S): CASREACT 142:355052; MARPAT 142:355052

Amidines ACH(CH3)C(:NR)NHR1 [A = (un)substituted Ph; benzoyl,
(un)substituted heteroaryl; R = H, Cl-5 alkyl, phenylalkyl, alkenyl,
cycloalkyl, alkoxy, etc.; R1 = H, Me, Et; e.g., (R, S)-2-(4isobutylphenyl)propionamidine hydrochloridel, useful in the inhibition of
chemotaxis of neutrophils induced by interleukin-8 in the treatment of
psoriasis, ulcerative colitis, melanoma, chronic obstructive pulmonary
disease, bullous pemphigo, rheumatoid arthritis, idiopathic fibrosis,
glomerulonephritis, and in the prevention and treatment of damages caused
by ischemia and reperfusion., are prepared
261178-48-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)

(in the preparation of amidines and their salts useful in the
sibition of

Page 10 10/568,760

ANSWER 10 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) chemotaxis of neutrophils induced by interleukin-8) 261178-48-7 CAPLUS

Benzeneethanimidamide, N-hydroxy-α-methyl-4-(2-methylpropyl)- (CA

849063-66-7P 849063-67-8P RL: PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of amidines and their salts useful in the inhibition of chemotaxis of neutrophils induced by interleukin-8) 849063-66-7 CAPLUS Benzeneethanimidamide, α-methyl-4-(2-methylpropyl)-, (+)- (CA INDEX NAME)

849063-67-8 CAPLUS Benzeneethanimidamide, α -methyl-4-(2-methylpropyl)-, (-)- (CA INDEX NAME)

Rotation (-).

849063-49-6P 849063-50-9P 849063-51-0P 849063-52-1P 849063-53-2P 849063-54-3P 849063-56-5P 849063-57-6P 849063-58-7P 849063-59-8P 849063-60-1P 849063-61-2P

ANSWER 10 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

849063-52-1 CAPLUS Benzeneethanimidamide, 3-benzoyl- α -methyl-, hydrochloride (1:1) (CA INDEX NAME)

849063-53-2 CAPLUS [1,1'-Biphenyl]-4-ethanimidamide, 2-fluoro- α -methyl-, hydrochloride (1:1) (CA INDEX NAME)

849063-54-3 CAPLUS Methanesulfonic acid, 1,1,1-trifluoro-, 4-(2-amino-2-imino-1-methylethyl)phenyl ester, hydrochloride (1:1) (CA INDEX NAME)

L4 ANSWER 10 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) RL: SFN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (USes) (prepn. of amidines and their salts useful in the inhibition of chemotaxis of neutrophils induced by interleukin-8) RN 84963-49-6 CAPLUS CN Benzenethanimidamide, α-methyl-4-(2-methylpropyl)-, hydrochloride (1:1) (CA INDEX NAME)

● HC1

849063-50-9 CAPLUS Benzeneethanimidamide, $\alpha\text{-methyl-}4\text{-}(2\text{-methylpropyl})\text{-}, hydrochloride (1:1), (+)- (CA INDEX NAME)$

HCl

849063-51-0 CAPLUS Benzeneethanimidamide, α -methyl-4-(2-methylpropyl)-, hydrochloride (1:1), (-)- (CA INDEX NAME)

Rotation (-).

L4 ANSWER 10 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

● HCl

849063-56-5 CAPLUS Benzeneethanimidamide, α -methyl-4-(2-methylpropyl)-N-[3-(1-piperidinyl)propyl]-, hydrochloride (1:2) (CA INDEX NAME)

●2 HC1

849063-57-6 CAPLUS Benzeneethanimidamide, N, α -dimethyl-4-(2-methylpropyl)-, hydrochloride (1:1) (CA INDEX NAME)

● HC1

849063-58-7 CAPLUS Benzeneethanimidamide, 3-benzoyl-N-[3-(dimethylamino)propyl]- α -methyl-, hydrochloride (1:2) (CA INDEX NAME)

10/568,760

L4 ANSWER 10 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

Page 11

NH- (CH2) 3-NMe2

●2 HC1

849063-59-8 CAPLUS Benzeneethanimidamide, α -methyl-4-(2-methylpropyl)-, acetate (1:1) (CA INDEX NAME)

CM 1

CRN 487007-20-5 CMF C13 H20 N2

CM

849063-60-1 CAPLUS Benzeneethanimidamide, N-[3-(dimethylamino)propyl]- α -methyl-4-(2-methyl-propyl)- (CA INDEX NAME)

L4 ANSWER 10 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

$$\begin{array}{c} \text{Me}_{2}\text{N-}\left(\text{CH}_{2}\right)_{3} - \text{NH-C-CH} \\ \text{HN Me} \end{array}$$

849063_61_2 CAPLUS

Benzeneethanimidamide, α-methyl-4-(2-methylpropyl)-N-(phenylmethyl)(CA INDEX NAME)

(Continued)

L4 ANSWER 11 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2005:150032 CAPLUS

DOCUMENT NUMBER: 142:411301

Thibition of secretory phospholipase A2. 2-Synthesis and structure-activity relationship studies of 4,5-dihydro-3-(4-tetradecyloxybenzy)-1,2,4-4-doxadiazol-5-one (PMS1062) derivatives specific for group II enzyme

AUTHOR(S): Dong, Chang-Zhi, Ahamada-Himidi, Azali; Plocki, Stephanie; Aoun, Darina; Touaibia, Mohamed;

Habich, Nadia; Huet, Jack; Redeuilh, Catherine;
Cmbetta, Jean-Edouard; Godfroid, Jean-Jacques;
Massicot, France; Heymans, Francoise
Unite de Pharmacochimie Moleculaire et Systemes
Membranaires (EA2381), Laboratoire de Pharmacochimie
Moleculaire, Universite Paris 7-Denis Diderot, Paris,
75251, Fr.
Bioorganic & Medicinal Chemistry (2005), 13(6),
1989-2007
CODEN: BMECEP; ISSN: 0968-0896
Elsewier Ltd.
Journal

SOURCE:

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
GI Journal

English CASREACT 142:411301

CORPORATE SOURCE:

The discovery of a series of specific inhibitors of human group IIA phospholipase A2 (hGIIA PLA2) displaying promising in vitro and in vivo properties has been recently reported. Here the influence of different structural modifications on the specificity and potency of oxadiazolones, e.g. I [X = CH2, CH2CH2, CHMe, CMe2; R = MeO, n-octyloxy, n-tetradecylthio, N,N-di(heptyl)amino, etc.], against hGIIA PLA2 vs. porcine group IB PLA2 is described. The SAR results, as well as the log

and pKa values of the oxadiazolones studied provide important information towards the comprehension of the mode of action of this kind of compds. 310869-86-4P 850143-48-5P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and calculated hydrophobicity of ether, thioether or amino-functionalized aralkyl oxadiazolones as inhibitors of human secretory phospholipase A2 specific for group II enzyme) 310869-86-4 CAPLUS Benzeneethanimidamide, N-hydroxy-α-methyl-4-(tetradecyloxy)- (CA INDEX NAME)

L4 ANSWER 11 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

850143-48-5 CAPLUS Benzeneethanimidamide, N-hydroxy- α , α -dimethyl-4-(tetradecyloxy)- (CA INDEX NAME)

Me- (CH2)13

THERE ARE 26 CITED REFERENCES AVAILABLE FOR REFERENCE COUNT:

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

06/05/2008 Habte

ANSWER 12 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 2004:550957 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 141:106464 TITLE: Preparation of pyrazolo[3,4-b]pyridine derivatives use in pharmaceutical compositions as phosphodiesterase inhibitors allen, David George; Coe, Diane Mary; Cook, Caroline Mary; Cooper, Anthony William James; Dowle, Michael Dennis; Edlin, Christopher David; Hamblin, Julie Nicole; Johnson, Martin Redpath; Jones, Paul Spencer; Lindwall, Mika Kristian; Mitchell, Charlotte Jane; Redgrave, Alison Judith Glaxo Group Limited, UK FCT Int. Appl., 244 pp. CODEN: PIXND2
Patent
English INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: DATE TG CA 2511340 A1 20040714 AU 2003-293999
A1 20040714 AU 2003-293999
EP 1581532 A1 20051005 EP 2003-789413 20031219
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
ER 2003017645 A 20051206 BR 2003-17645 20031219
CN 1751042 A 20060322 CN 2003-80109835 20031219
JP 2006513258 T 20060420 JP 2005-502565 20031219
JP 2006513258 T 20050630 AU 2004-295277 20041217
CA 20040257004 20041217
CA 2004-2557004 20041217 CA 2557004 WO 2005058 A1 A1 20050630 20050630 CA 2004-2557004 WO 2004-EP14490 2005058892 2004121 05892 Al 20050630 W0 2004-EP14490
AE, AG, AL, AM, AT, AL, AZ, BA, BB, BG, BR, BW,
CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG,
GE, GH, GW, HR, HU, ID, IL, IN, IS, JP, RE, KG,
LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MM, MN,
NO, NZ, CW, PG, PH, PL, PT, RO, RU, SC, SD, SE,
TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN,
BW, GH, GW, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ,
AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH,
EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, BY, BZ, CA, CH, ES, FI, GB, GD, KP, KR, KZ, LC, MX, MZ, NA, NI, w: NI, SY, ZW MX, SG, YU, UG, CY, MC, SK, ZA, ZM, CZ, NL, SL, ZM, RW:

L4 ANSWER 12 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

RC, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GN, ML,

MR, NE, SN, TD, TG

EP 1737857 A1 20070103 EP 2004-804089 20041217 CN 1914205 JP 2007514704 ZA 2005005074 IN 2005KN01207 MX 2005PA06923 NO 2005003600 20070607 20060927 20070713 20050818 20050822 20061109 20070517 20070518 ZA 2005-5074
IN 2005-KN1207
MX 2005-PA6923
NO 2005-3600
US 2006-540371
US 2006-596561
IN 2006-KN1988
NO 2006-3340
GB 2002-30045 20050622 20050623 20050722 US 20060252790
US 20060252790
US 20070111995
IN 2006KN01988
NO 2006003340
PRIORITY APPLN. INFO.: 20050722 20060221 20060616 20060714 20060718 GB 2002-30165 A 20021224 GB 2003-7998 A 20030407 W 20031219 WO 2003-EP14867 WO 2003-EP314867 A 20040316 A 20040325 GB 2004-6754 WO 2004-EP14490 W 20041217

MARPAT 141:106464

ANSWER 13 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

2003:235774 CAPLUS

OTHER SOURCE(S):

ACCESSION NUMBER:

ANSWER 12 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

Pyrazolo[3,4-b]pyridine derivs., such as I [R = heterocyclyl; Rl = (CH2)2OH, alkyl, fluoroalkyl; R2 = H, Me, fluoroalkyl; R3 = alkyl, (un)substituted-Ph, cycloalkyl, heterocyclyl, etc.; R3a = H, alkyl, were prepared for therapeutic uses as inhibitors of phosphodiesterase, particularly phosphodiesterase IV (PDE4). These pyrazolo[3,4-b]pyridines were claimed for use in the treatment and/or prophylaxis of cognitive impairment and inflammatory and/or allergic diseases, such as chronic obstructive pulmonary disease (COPD), asthma, or allergic rhinitis.

Thus.

pyrazolo[3,4-b]pyridine derivative II was prepared via a cvclc

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of pyrazolo[3,4-b]pyridine derivs. for use in pharmaceutical

compns. as phosphodiesterase inhibitors)
42191-51-5 CAPLUS
Benzeneethanimidamide, N-hydroxy-α-methyl- (CA INDEX NAME)

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT:

L4 ANSWER 13 OF 60 CAPUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 2003:225774 CAPLUS
DOCUMENT NUMBER: 138:368454
TITLE: Photochemistry of Crystalline Chlorodiazirines: The
Influence of Conformational Disorder and
Intermolecular C1···NN
Interactions on the Solid-State Reactivity of Singlet
Chlorocarbenes
AUTHOR(S): Sanrame, Carlos N.; Suhrada, Christopher P.; Dang,
Bung; Garcia-Garibay, Miguel A.

CORPORATE SOURCE: Department of Chemistry and Blochemistry, University
of California, Los Angeles, CA, 90095-1569, USA
JOURNAI of Physical Chemistry A (2003), 107(18),
3287-3294
COEDE: JPCAFH; ISSN: 1089-5639
American Chemical Society
JOURNAID TYPE:
LANGUAGE: English
CTHER SOURCE(S): CASEACT 138:368454
AB A photochem. study was carried out with 3-R-substituted
3-chlorodiazirines
with 4-biphenyl- (4a), (4-biphenyl)methyl (4d) substituents. The
chlorodiazirines were prepared from the corresponding amidines by a
procedure involving oxidation with tert-Bu hypochlorite under
catalvais. The crystalline nature of 4ard was established by catalysis. The crystalline nature of 4a-d was established by differential erential scanning calorimetric anal., which revealed melting endotherms prior to thermal decomposition Photochem. results in crystalline solids were those observed in solution, and the products were analyzed in terms of corresponding singlet-state chlorocarbene intermediates (5a-d). Irradiation of 4a in solution and in crystals resulted in formation of azine RCIC:NN:CCIR NNN:CCIR

9a (R = C6H4-p-Ph) by reaction of carbene 5a with its precursor. Equally selective, diazirine 4d gave alkene Me2C:CCl(C6H4-p-Ph) 6d as the only product by a 1,2-Ph migration from carbene 5d. In contrast, irradiation compds. 4b and 4c resulted in formation of two alkenes by 1,2-H shifts formation of azines by reactions of the carbenes with their precursors. The low selectivity of 4D was rationalized in terms of structural data from single-crystal X-ray diffraction anal., which revealed two dered disordered

diazirine conformers and close Cl...N contacts

between adjacent mols. Rapid conformational equilibration in the solid
state was also suggested by solid-state 13C CPMAS NMR. Similar
structural

effects are also postulated to account for the solid-state reactivity of 524068-7/-7

RE: RCT (Reactant); RACT (Reactant or reagent)

(PTC oxidation; the influence of conformational disorder and intermol.

Cl...N:N interactions on the solid-state
reactivity of singlet chlorocarbenes formed in photolysis of

3-chlorodiazirines)

06/05/2008 Habt.e

Page 13 10/568,760

ANSWER 13 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN 524068-77-7 CAPLUS (Continued) RN

[1,1'-Biphenyl]-4-ethanimidamide, α,α -dimethyl-, hydrochloride (1:1) (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 74 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

ANSWER 14 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

THERE ARE 16 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

SOURCE:

IT 374063-57-7P
RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (synthesis and activity of imidoyl thioureas as non-nucleoside reverse transcriptase inhibitors)
RN 374063-57-7 CAPLUS
CN Benzeneethanimidamide, 2,6-dichloron-F[[(4-cyanophenyl)amino]thioxomethyl]α-methyl- (CA INDEX NAME) ANSWER 15 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN PLUS COPYRIGHT 2008 ACS on STN 2001:265374 CAPLUS 2001:265374 CAPLUS 134:280609 Preparation of N-{a- (cyclopropylmethoximino)aralkyl]phenylacetamides and analogs as agrochemical fungicides Rheinheimer, Joachim; Eicken, Karl; Rose, Ingo; ACCESSION NUMBER: DOCUMENT NUMBER: INVENTOR(S):

Stemming from work on a previous clin. candidate, loviride, and other α -APA derivs., a new series of potent non-nucleoside reverse transcriptase inhibitors (NNRTIs) has been synthesized. The ITU analogs, which contain a unique diarylated imidoyl thiourea, e.g. (I), are very active in inhibiting both wild-type and clin. important mutant strains of

ANSWER 14 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

135:371498

13513/1498 Evolution of anti-HIV drug candidates. Part 1: From α -Anilinophenylacetamide (α -APA) to imidoyl thiourea (ITU) Ludovici, D. W.; Kukla, M. J.; Grous, P. G.;

S.; Andries, K.; de Bethune, M.-P.; Azijn, H.; Pauwels, R.; De Clercq, E.; Arnold, E.; Janssen, P.

Janssen Research Foundation, Spring House, PA, 19477,

USA Bioorganic & Medicinal Chemistry Letters (2001), 11(17), 2225-2228 CODEN: BMCLE8; ISSN: 0960-894X Elsevier Science Ltd. Journal English CASREACT 135:371498

ACCESSION NUMBER:

DOCUMENT NUMBER:

CORPORATE SOURCE:

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
GI

HIV-1. 374063-57-7P

TITLE: AUTHOR(S):

SOURCE:

AB

IT

Thomas; Ammermann, Eberhard; Speakman, John-Bryan; Strathmann, Siegfried; Lorenz, Gisela BASF A.-G., Germany PCT Int. Appl., 34 pp. CODEN: PIXXD2 Patent LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. APPLICATION NO. KIND DATE DATE WO 2001025187 A2 A3 20010412 20011101 WO 2000-EP9744 20001005 US 2005-61470 US 20050187265 20050825 20050222 US 7101900 20060905 PRIORITY APPLN. INFO.: A 19991006 DE 1999-19948266 WO 2000-EP9744 W 20001005

OTHER SOURCE(S): MARPAT 134:280609 CTHER SOURCE(S): MARPAT 134;280609

AB R1ZC(:NOR)NHCOR2 (R = cyclopropylmethyl)[I, Rl = (un)substituted Ph, -pyridyl, -thienyl; R2 = (un)substituted phenyl-, -thienyl-, -pyrazolylalkyl; Z = (un)substituted (heteroatom- or cyclopropylene-interrupted) alkylene] were prepared Thus, BONH2 was added to 2,6-c12c6H3CH2CN and the resulting amidoxime O-alkylated by cyclopropylmethyl bromide to give, after N-acylation, I (Rl = 2,6-c12c6H3, R2 = CH2Ph, Z = CH2). Data for biol. activity of I were given.

IT 333748-79-1P

US 2002-89148

A3 20020327

ANSWER 15 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)
RL: AGR (Agricultural use); BAC (Biological activity or effector, except
adverse); BSU (Biological study, unclassified); SPN (Synthetic
preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)
(prepn. of N-[a-(cyclopropylmethoximino)aralkyl]phenylacetamides
and analogs as agrochem. fungicides)
333748-79-1 CAPLUS
Benzeneacetamide, N-[1-[(cyclopropylmethoxy)amino]-2-phenylpropylidene](9CI) (CA INDEX NAME)

42191-51-5P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation of N-[α-(cyclopropylmethoximino)aralkyl]phenylacetamides and analogs as agrochem. fungicides)
42191-51-5 CAPLUS
Benzeneethanimidamide, N-hydroxy-α-methyl- (CA INDEX NAME)

ANSWER 16 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN EP 1999-126035 (Continued) A 19991227

> WO 2000-EP7358 W 20000731

OTHER SOURCE(S): MARPAT 134:178571

The title compds. (I) [p = 0-4; X = 0, S, NR5, or a direct bond; or XR2 taken together = CN; R1 = independently C(O)2R14, (un)substituted alkyl, halo, OH, SH, alkoxy, alkylthio, alkylcarbonyloxy, aryl, CN, NO2, hetercyclyl, R6, or NR7R8; R2 = hetercoyclyl, (un)substituted cycloalkyl, alkoxy, or alkylthio, heterocyclyl(oxy), heterocyclylthio, etc.; R3 and

independently H or (cyclo)alkyl; or R3 and R4 taken together form an alkenediyl; R5 = H or alkyl; R6 = (un)substituted (cyclo)alkylsulfonyl, amino(alkylsulfonyl, heterocyclylsulfonyl, etc.; R7 and R8 = independently H, (cyclo)alkyl, (di)hydroxyalkyl, mercaptoalkyl, aryl(alkyl), alkyloxyalkyl, alkyl(thio)carbonyl, aryl(thio)carbonyl, heterocyclyl(thio)carbonyl, (O)ZR14, or (un)substituted aminocarbonyl, etc.; or R7 and R8 together with the N to which they are attached form pyrrolinone, piperidinone, or hexahydroazepinone; R14 = H, alkynyl, or (un)substituted (alkyl)acyl, alkyl, alkenyl, heterocyclyl, etc.; Z = O,

NH, CH2O, or CH2S; or ZR14 taken together = CH2CN or CH2PO3H2 and its esters] and their N-oxides, pharmaceutically acceptable salts, or stereochem. isomers were prepared as selective chemokine inhibitors. Fexample, 2,6-dichlor-4-(4,5-dihydro-3,5-dioxo-1,2,4-triazin-2(3H)-yl)-a,a-dimethylbenzeneethanethioamide was coupled with Et β -bromo-y-oxobenzenebutanoate (46.5%), cyclized to form the thiazoleacetic acid (79%), and esterified with 3-bromodihydro-2(3H)-furanone to give II. As selective interleukin 5 (IL-5) and monocyte

ANSWER 16 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN 2001:115148 CAPLUS

DOCUMENT NUMBER: 134:178571

TITLE:

134:178571
Preparation of 6-azauracil derivatives as interleukin-5 inhibitors
Lacrampe, Jean Fernand Armand; Freyne, Eddy Jean Edgard; Deroose, Frederik Dirk; Fortin, Jerome Michel Claude; Coesemans, Erwin
Janssen Plarmaceutica N.V., Belg.
PCT Int. Appl., 163 pp.
CODEN: PIXXD2
Patent INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: English

PAT	TENT	NO.			KIN												ATE	
wo	2001	0108	 66					0215									0000	
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB	, во	, в	R,	BY,	BZ,	CA,	CH,	CN.
		CR.	CU.	CZ.	DE.	DK.	DM.	DZ.	EE.	ES	. FI	, G	В.	GD,	GE,	GH.	GM.	HR
		HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KF	, KF	, K	z,	LC,	LK,	LR,	LS,	LT.
		LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX	, M2	, N	ю,	NZ,	PL,	PT,	RO,	RU
		SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TF	, TI	, т	z,	UA,	UG,	US,	UZ,	VN
		YU,	ZA.	ZW														
	RW:	GH.	GM.	KE.	LS.	MW.	MZ.	SD,	SL.	SZ	. T2	. U	ſĠ,	ZW.	AT.	BE.	CH.	CY.
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	II	, LU	, M	ic,	NL,	PT,	SE,	BF,	BJ
		CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MB	, NE	, s	N,	TD,	TG			
CA	2380	759			A1		2001	0215		CA	2000	-23	80.	759		2	0000	731
BR	2000	0130	14		A		2002	0416		BR	2000	-13	01	4		2	0000	731
EP	2000 1206	471			A1		2002	0522		EP	2000	-94	80:	15		2	0000	731
EP	1206	471			В1		2006	0301										
	R:	AT,						FR,		GP	, II	, L	ı,	LU,	NL,	SE,	MC,	PT.
		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL								
TR	200200310 7 2002002692 7 2002002692 9 2003506451				T2		2002	0821		TR	2002	-31	0			2	0000	731
HU	2002	0026	92		A2		2002	1228		HU	2002	-26	92			2	0000	731
HU	2002	0026	92		A3		2003	0128										
JP	2003	5064	51		Т		2003	0218		JP	2001	-51	56	75		2	0000	731
	2002									EE	2002	-57				2	0000	731
	5165						2004	0227						06				
	7800				B2			0224		ΑU	2000	-61	609	9		2	0000	731
	3188				Т			0315		AΤ	2000	-94	80:	15 15		2	0000	731
	2260				Т3		2006			ES	2000	-94	80:	15		2	0000	
	2714				В		2007							5824				
	7954						2008							0.4				
	1063						2002							57				
IN	2002	MN00	144		A		2005	0318		IN	2002	-MN	114	4		2	0020	131
NO	2002 3223	0005	65		A		2002	0326		NO	2002	-56	5			2	0020	205
NO	3223	86			B1		2006	0925										
za	2002	0010	07		A		2003	0505										
	2002													43				
	2003		453		A1					US	2002	-75	876	5		2	0020	214
	6911				B2		2005	0628										
	1048				A1		2005	0930		HK	2003	-10	07:	18		2	0030	128
IT	/ APP	LN.	INFO	. :						EΡ	1999	-87	01	70		A 1	9990	806

ANSWER 16 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) chemotactic protein-1 and -3 (MCP-1 and MCP-3) inhibitors, I are useful for treating eosinophil-dependent inflammatory diseases, esp. bronchial asthma (no data). Processes using I for marking receptors and imaging organs via radiolabeling are also claimed. 261512-63-49 25968-68-1P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (intermediate; preparation of IL-5 inhibiting thiazolylalkylphenyl-6-azauracil derivs. by coupling of 4-dioxotriazinyl-a,a-dimethylbenzeneethanethioanides with a-oxoalkyl halides, cyclization, and addition of functionally substituted groups) 261512-63-4 CAPLUS
Benzeneethanimidamide, 4-amino-2,6-dichloro-N-hydroxy-a,a-dimethyl- (CA INDEX NAME)

RN CN

325968-68-1 CAPLUS Benzamide, N-[2-(4-amino-2,6-dichloropheny1)-1-imino-2-methylpropy1]-2-methyl- (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

06/05/2008 Habte

ANSWER 17 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 2001:63983 CAPLUS MENT NUMBER: 134:131527 ACCESSION NUMBER:

DOCUMENT NUMBER:

134:131527
Preparation and effect of heteroaromatic ring compounds against autoimmune disorders and chronic inflammation
Nakatsuka, Masashi; Nakatani, Shogo; Okada,
Shin-ichiro; Tsuboi, Katsunori; Nishikaku, Fumio Sumitomo Pharmaceuticals Co., Ltd., Japan PCT Int. Appl., 190 pp.
CCDEN: PIXXD2
Patent TITLE:

INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE: Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT	NO.			KIN	D	DATE			APPL	ICAT	ION :	NO.		D.	ATE	
					_									-		
WO 2001005774			A1		2001	0125		WO 2	000-	JP46	16		2	0000	710	
W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,
	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,
	YU,	ZA,	ZW													
RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	BJ,

DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, FT, SE, BF, BJ, CF, GG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
CA 2377527 A1 20010125 CA 2000-2377527 20000710
F1 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL
PRIORITY APPLN. INFO::

JP 1999-201447 A 19990715

JP 2000-58217 A 20000303

WO 2000-JP4616 W 20000710

OTHER SOURCE(S): MARPAT 134:131527 L4 ANSWER 17 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

Title compds. [I; R1 = F, C6H5CO, C6H5CHO2(CH2)2; R2 = H, C6H5; R3 = H, CH3; R4 = H, CH3; R5 = CH3, CH2CH2N[(CH2)2]O; R6 = H, CH3; R4-R5 = CH2CH2OCH2CH2, CH2CH2SCH2CH2, CH2CH2S(H0)(100)CH2CH2; R6 = H, CH3; R7 = CH3, H, CH2CH2OH, CN, C(MHN) ((CH2)2]O; R5-R7 = CH2CH2C, CH2CH2CH2, CH2CH0CH2; R6-R7 = CH2CH2OCH2CH2; R8 = H, CH3; R9 = H, CH3; X = N, NCH3, S; Y = NCH3, S, NI, NSO2C6H5; Z = CH, O, S, N; dotted line = single, double bond] and pharmaceutically acceptable salts exhibiting excellent phys. properties and potent ameliorative effects against both immune disorders and chronic inflammation are prepared. Thus, the title ound II compound II

ound II

was prepared and tested.
321879-91-8P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(preparation and effect of heteroarom, ring compds. against immune

disorders

cders and chronic inflammation)
321879-91-8 CAPLUS
[1,1'-Biphenyl]-4-ethanimidamide, N-chloro-2-fluoro-α-methyl- (CA
INDEX NAME)

L4 ANSWER 17 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

Habte

ANSMER 18 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 2000:900841 CAPLUS 134:37031 E: FVIIA/TF activity inhibiting compounds Jakobsen, Palle; Persson, Egon Nova Nordisk A/s, Den. CE: CE: CODEN: FIXXD2 PAtent UNGGE: English LY ACC. NUM. COUNT: 4 L4 ANSWER 18 OF 6(
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE: DOCUMENT TYPE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PATENT NO.					KIN		DATE										
T/s	10	2000	0772	46		A2		2000 2001	1221		WO.	2000-	DK31	6			20000	613
			AE, CU, ID, LV,	AG, CZ, IL, MA, SG,	AL, DE, IN, MD,	AM, DK, IS, MG,	AT, DM, JP, MK,	AU, DZ, KE, MN,	AZ, EE, KG, MW,	BA, ES, KP, MX,	FI KF M2	B, BG, GB, K, KZ, I, NO, T, TZ,	GD, LC, NZ,	GE, LK, PL,	GH, LR, PT,	GM LS RO	HR, LT, RU,	HU, LU, SD,
			GH, DE, CF,	GM, DK, CG,	ES,	FI,	FR, GA,	GB,	GR, GW,	IE, ML,	IT MF	, TZ, , LU,	MC, SN,	NL, TD,	PT, TG	SE	BF,	вЈ,
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GF	2000- , IT,	LI,	LU,	NL,	SE	, MC,	PT,
	IS	6444	5308: 878 434	19		T B1 B1		2003 2001 2002	0903		US	2001- 2000- 2001-	8448	28			20010	427
PRIORI						Al		2003	0417		DK	2002- 1999-	840			A		614
												1999- 1999-						
												1999-						
												1999-						
												2000-						
												2000-						
											US	2000-	6160	10			20000	

The invention relates to compds. inhibiting the activation of FX to FXa

by

TF/FVIIa. The compds. are anticoagulants. The invention also relates to a method of identifying a drug candidate.

IT 313236-51-0
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study);

Page 16 10/568,760

ANSWER 18 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) (FVIIA/TF activity inhibiting compds.) 236-51-0 CAPLUS

Senzeneethanimidamide, α -methyl-4-(2-methylpropyl)-N-[2-[4-(2-methylpropyl)phenyl]propyl]- (CA INDEX NAME)

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. R SOURCE(S): MARPAT 134:25351
The invention provides phospholipase A2 inhibitor heterocyclic compds. (Markush included). The compds. are capable of acting on PLA2 and are advantageously secreted nonpancreatic PLA2-specific inhibiting compds. completely inactive towards pancreatic PLA2. The invention also provides a method for preparing the compds., pharmaceutical and cosmetic compns. containing them, and their use in particular for treating inflammatory OTHER SOURCE(S):

134:25351

Heterocyclic phospholipase A2-specific inhibitors, their preparation, their use in treatment of inflammation, and pharmaceutical and cosmetic compositions containing them Assogba, Leon; Heymans, Francoise; Dong, Chang-Zhi; Godfroid, Jean-Jacques Universite Paris 7 - Denis Diderot, Fr. PCT Int. Appl., 64 pp.
CODEN: PIXXD2

ANSWER 19 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

134:25351

pathologies. 310869-86-4P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction; heterocyclic phospholipase A2-specific inhibitor

preparation, use in treatment of inflammation, and pharmaceutical and
 cosmetic compns. containing them)
310869-86-4 CAPLUS
Benzeneethanimidamide, N-hydroxy-α-methyl-4-(tetradecyloxy)- (CA

DOCUMENT NUMBER:

PATENT ASSIGNEE(S):

INVENTOR(S):

TITLE:

INDEX NAME)

L4 ANSWER 19 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

THERE ARE 18 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

ANSWER 20 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2000:190770 CAPLUS
132:222555
Preparation of interleukin-5 inhibiting 6-azauracil derivatives
Freyne, Eddy Jean Edgard; Lacrampe, Jean Fernand Armand; Deroose, Frederik Dirk; Venet, Marc Gaston Janssen Pharmaceutica N.V., Belg.
Eur. Pat. Appl., 37 pp.
CODEN: EPEXDW
Patent
English
1 2000:190770 DOCUMENT NUMBER: INVENTOR(S): PATENT ASSIGNEE(S): DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE EF 1114046 B1 20030423 E: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
JP 2002526495 T 20020820 JP 2000-574104 19990914 JP 2000-574104 AT 1999-947336 ES 1999-947336 US 2001-812731 AT 238301 ES 2198958 US 20020010177 US 6894046 20030515 20040201 19990914 20020124 20010319 В2 20050517 PRIORITY APPLN. INFO.: EP 1998-203148 A 19980918 WO 1999-EP6776 W 19990914

OTHER SOURCE(S): MARPAT 132:222555

06/05/2008 Habt.e

L4 ANSWER 20 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

The title compds. [I; p = 0-4; X = 0, S, NR5, a direct bond; Y = 0, S, NR5, SO2; R1 = alkyl, halo, polyhaloalkyl, etc.; R2 = Hetl, cycloalkyl, alkyl, and if X = 0, S, NR5, then R2 may also represent aminocarbonyl, alkylcarbonyl, etc.; R3, R4 = H, alkyl, cycloalkyl; R3R4 = alkanedkyl; R5 = H, alkyl; Hetl = (un)substituted heterocyclel, useful for treating eosinophil-dependent inflammatory diseases, and marking a receptor, were prepared and formulated. E.g., a multi-step synthesis of 1,2,4-triazine-3,5(2H,4H)-dione II which showed 90.5% inhibition of II.-5 production, was given. 261512-63-49 261512-64-5P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation of interleukin-5 inhibiting 6-azauracil derivs.) 261512-63-4 CAPLUS
Benzeneethanimidamide, 4-amino-2,6-dichloro-N-hydroxy-α,α-dimethyl- (CA INDEX NAME)

261512-64-5 CAPLUS Benzeneethanimidamide, 4-amino-2,6-dichloro-N-[(2-chlorobenzoyl)oxy]- α , α -dimethyl- (9CI) (CA INDEX NAME)

ACCESSION NUMBER:

ANSWER 21 OF 60 CAPLUS COPYRIGHT 2008 ACS ON STN

SSSION NUMBER: 2000:146378 CAPLUS

MENT NUMBER: 132:293711

SORATE SOURCE: Geffken, Detlef; Holst, Carsten; Willrodt, Inke

COPARTE SOURCE: Institute of Pharmacy, Department of Pharmaceutical

Chemistry, University of Hamburg, Hamburg, 20146,

Germany

MEL: CODEN: HCOMEX; ISSN: 0793-0283

ISHER: Feund Publishing House Ltd.

MINDIT TYPE: Journal

SUAGE: English

CASREACT 132:293711 DOCUMENT NUMBER: TITLE: AUTHOR(S): CORPORATE SOURCE:

SOURCE:

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
GI

Treatment of 4-hydrazono-2-oxazolidinones with acetic anhydride afforded novel 2-acetyloxazolo[4,3-c]-1,2,4-triazol-5-ones (I; R1 = Me, H; R2 = $\frac{1}{2}$ AB

Me,

Ph, H, 4-fluorophenyl; CR2R3 = CHPh, cyclopentylidene, CMe2, CPh2, tetrahydropyran-4-ylidene, tetrahydrothiopyran-4-ylidene) in good yields. 264124-05-2P 264124-09-6P RE. RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and cyclic carbonylation of) 264124-05-2 CAPLUS Benzeneethanimidic acid, α-hydroxy-α-methyl-, (tetrahydro-4H-pyran-4-ylidene)hydrazide (9CI) (CA INDEX NAME) IT

264124-09-6 CAPLUS Benzeneethanimidic acid, 4-fluoro- α -hydroxy- α -methyl-, (tetrahydro-4H-thiopyran-4-ylidene)hydrazide (9CI) (CA INDEX NAME)

Habte

ANSWER 20 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)
RENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT:

FORMAT

ANSWER 21 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

160154-90-5 RL: RCT (Reactant); RACT (Reactant or reagent) (reaction with tetrahydropyranone) 160154-90-5 CAPLUS Benzeneethanimidic acid, α -hydroxy- α -methyl-, hydrazide (CA INDEX NAME)

264124-07-4 RL: RCT (Reactant); RACT (Reactant or reagent) (reaction with tetrahydrothiopyranone) 264124-07-4 CAPLUS Benzeneethanimidic acid, 4-fluoro- α -hydroxy- α -methyl-, hydrazide (CA INDEX NAME)

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE REFERENCE COUNT:

FORMAT

06/05/2008

ANSWER 22 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1999:27808 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 130:81527 Preparation of novel amidrazone derivatives having TITLE: Preparation of novel amidrazone derivatives having antifungal activity Kageyama, Shunji; Kontani, Toru; Fujii, Masahiro; Igarashi, Kiyoshi; Yamamoto, Osamu Yamanouchi Pharmaceutical Co., Ltd., Japan PCT Int. Appl., 48 pp. CODEN: PIXND2 INVENTOR(S): PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE WO 9858905 Al 19981230 WO 1998-JP2817 19980624
Wi AL, AM, AU, AZ, BA, BB, BG, BR, BY, CA, CN, CU, CZ, EE, GE, GH,
GW, HU, ID, IL, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LV,
MD, MG, MK, MN, MW, MX, NO, NZ, PL, RO, RU, SD, SG, SI, SK, SL,
TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW
RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES,
FI, FR, GB, GR, IE, IT, LU, MC, NL, FT, SE, BF, BJ, CF, CG, CI,
CM, GA, GN, ML, MR, NE, SN, TD, TG

AU 9879330 A 19980624
PRIORITY APPLN: INFO:: JP 1997-168354 A 19970625

WO 1998-JP2817 W 19980624

OTHER SOURCE(S): MARPAT 130:81527

Amidrazone derivs. of formula [I, wherein the ring Ra represents: (1) an optionally substituted monocyclic to tricyclic aromatic hydrocarbon, (2) AB

optionally substituted monocyclic to tricyclic saturated or unsatd.

ro ring containing one or more hetero atoms selected from N, O and S, (3) an optionally substituted and optionally cross-linked cycloalkyl, or (4) an optionally substituted and optionally cross-linked cycloalkenyl; the ring Rb represents (1) an optionally substituted monocyclic to tricyclic

hydrocarbon or (2) an optionally substituted monocyclic to tricyclic saturated

L4 ANSWER 22 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

218918-70-8 CAPLUS Benzeneethanimidic acid, 2,4-difluoro- α -hydroxy- α -methyl-, 2-(4-chlorophenyl)hydrazide, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 22 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) ANSWER 22 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) or unsatd. hetero ring contg. one or more hetero atoms selected from N, O and S; one of Rc and Rd represents H and the other is not present; Re represents H or OH; Rf represents H or lower alkyl, or YRal; the dotted line "..." represents a single bond or a double bond; n is 1 to 8; A represents a bond or a lower alkylene optionally substituted by a lower alkyl; and X represents a bond, CO, CO2, CONRg, COCONRG1, CH:CHCONRG2, NRg3, NRg4CO, NRg5CO2, NRg5CONRG7, O, O2C, O2CNRG8, OCH2CONRG\$, S, SO, SO2, SO2NRG10, or SO2NRG11CO; wherein Rg and Rg1 - Rg11 represent H,

alkyl, or YRa2; Ra1 and Ra2 represents the same group as Ra; Y represents a single bond, CH2, or CO; a proviso given] or pharmaceutically

alkyl, or YMa2; Mai and Ma2 represents the same group as Ma; Y represents a single bond, CH2, or CO; a proviso given] or pharmaceutically acceptable salts thereof are prepd. Also claimed are pharmaceutical compns. thereof and a method for prevention or treatment of fungal or deep fungal infection by administration of I. These compds. I are useful for the treatment or prevention of fungal infection, in particular, deep fungal infection attributable to fungi, such as Candida, Aspergillus, and Cryptococcus. Thus, 2-(2-chloro-5-fluoro-6-oxo-1,6-dihydropyrimidin-1-yl)acetonitrile was treated with EtoH and HCI(g) in CHCI3 at 5° for 2 days to give a crude imidate which was condensed with 4-chlorophenylhydrazine hydrochloride in EtoH in the presence of EtoNa at room temp. overnight to give the title compd., 2-pyrimidinyl-N-phenylacetamidrazone (II). II showed 80% min. inhibitory concn. of 0.31, 0.31, and 0.63 mg/ml. against Candida albicans TIMM1768, Cryptococcus neoformans TIMM0362, and Aspergillus fumigatus TIMM1776, resp.

IT 218918-69-5P 218918-70-8F
RL: BAC (Biological activity or effector, except adverse); BSU (Biological)

(Biological

logical study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (USes) (preparation of novel amidrazone derivs. having antifungal activity) 218918-69-5 CAPLUS Benzeneethanimidic acid, 2,4-difluoro-a-hydroxy-a-methyl-, 2-phenylhydrazide, ethanedioate (1:1) (salt) (9CI) (CA INDEX NAME)

CRN 218918-68-4 CMF C15 H15 F2 N3 O

CM 2

CRN 144-62-7 CMF C2 H2 O4

ACCESSION NUMBER:

ANSWER 23 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
SSION NUMBER: 1998:573216 CAPLUS
MENT NUMBER: 129:196031
E: Racemic 2-Hydroxy-2-phenylpropanamidinium chloride DOCUMENT NUMBER: TITLE: and

(S)-2-Hydroxy-2-phenylbutanamidinium
(R)-2-Hydroxy-2-phenylethanoate
Barnes, John C.; Weakley, Timothy J. R.
Dep. Chem, Univ. Dundee, Dundee, Dul 4HN, UK
Acta Crystallographica, Section C: Crystal Structure
Communications (1998), C54(8), 1170-1173
CODEN: ACSCEE; ISSN: 0108-2701
Munksgaard International Publishers Ltd.
Journal

AUTHOR(S): CORPORATE SOURCE: SOURCE:

PUBLISHER: Munksgaard International Publishers ...

DOCUMENT TYPE: Journal
LANKOUAGE: Brglish
AB In 2-hydroxy-2-phenylpropanamidinium chloride, C9H13-N2O+.Cl-, the anion plays a central role in the H-bond network, chelating to one amidinium group and forming intermol. links to neighboring NH2 and OH- groups. The central feature in (S)-2-hydroxy-2-phenylbutanamidinium (R)-2-hydroxy-2-phenylethanoate, C10H15N2O+.C8H7O3-, is a ring linking the

and inter-mol. H bonds. Crystallog. data are given.

92442-87-0 RL: PRP (Properties)

(crystal structure of) 92442-87-0 CAPBUS Benzeneethanimidan.ide, α -hydroxy- α -methyl-, monohydrochloride (9CI) (CA INDEX NAME)

■ HC1

REFERENCE COUNT: 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

06/05/2008 Habt.e

ANSWER 24 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1998:294826 CAPLUS

ACCESSION NUMBER:

DOCUMENT NUMBER: 129:13472

TITLE: Amidrazones: a new class of coleopteran insecticide. Furch, J. A.; Kuhn, D. G.; Hunt, David A.; Asselin, M.; Baffic, S. P.; Diehl, R. E.; Palmer, Y. L.; Trotto, S. H.; Cyanamid Agric. Res. Cent., Am. Cyanamid Corp., Princeton, NJ, 08543-0400, USA AGS Symposium Series (1998), 686 (Synthesis and Chemistry of Agrochemicals V), 178-184 CODEN: ACSMC9; ISSN: 0937-6156 American Chemical Society Journal English Amidrazones: a new class of coleopteran insecticides AUTHOR(S):

CORPORATE SOURCE:

SOURCE

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

Amidrazones I (W = H, Cl, Br; X = H, CF3; Y = H, NO2, CF3; Z = H, Cl, Br, CF3, NO2) and related compds. were developed as insecticides specific against Coleoptera, especially Diabrotica undecimpunctata, with low AB toxicity to

Lepidoptera, Acarina, fish, birds and mice. The synthesis of the compds.

is outlined. 156820-05-2 156820-27-8

156820-05-2 156820-27-8
RL: AGR (Agricultural use); BIOL (Biological study); USES (Uses) (insecticide specific against Coleoptera)
156820-05-2 CAPLUS
Benzeneethanimidic acid, N-ethyl-α,α-dimethyl-,
2-[2,6-dichloro-4-(trifluoromethyl)phenyl]hydrazide (CA INDEX NAME)

156820-27-8 CAPLUS

196820-2/-6 CAPLOS Benzeneethanimidic acid, 4-chloro-N-ethyl-α,α-dimethyl-, 2-[2,6-dichloro-4-(trifluoromethyl)phenyl]hydrazide (CA INDEX NAME)

ANSWER 25 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: OCUMENT NUMBER:

PLUS COPYRIGHT 2008 ACS on STN
1998:112229 CAPLUS
128:192667
Preparation of substituted aromatic compounds as inhibitors of tumor necrosis factor and cyclic AMP phosphodiesterase
He, Wei; Hulme, Christopher; Huang, Fu-chih; Djuric, Stevan W.; Moriarty, Kevin; Labaudiniere, Richard Rhone-Poulenc Rorer Pharmaceuticals Inc., USA; He, Wei; Hulme, Christopher; Huang, Fu-Chih; Djuric, Stevan W.; Moriarty, Kevin; Labaudiniere, Richard PCT Int. Appl., 154 pp.
CODEN: PIXXD2
Patent
English
1 INVENTOR(S):

PATENT ASSIGNEE(S):

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

Habte

PATENT NO. APPLICATION NO. KIND DATE DATE A1 WO 9805327 19980212 WO 1997-US13343 19970722 9805327 A1 19980212 W0 1997-US13343 19970722
W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MN, MX, NO, NZ, PL, FT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN
RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GN, GR, IE, IT, LU, MC, NL, FT, SE, BF, BJ, CF, CG, CI, CM, GA, CM, ML, MR, NE, SN, TD, TG
9738990 A 19980225 A1 1998-23165 P 19960805 AU 1997-38990 US 1996-23165P 19970722 P 19960805 PRIORITY APPLN. INFO.: WO 1997-US13343 W 19970722

OTHER SOURCE(S): MARPAT 128:192667

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

This invention is directed to compound of formula [I; ring A = Q10, Q11;

This invention is directed to compound of formula [1; ring A = Q10, Q11;

= Q12, Q13, Q14; ring Ar2 = (un)substituted fused Ph or fused monocyclic heteroaryl; R = (un)substituted alkyl, aralkyl, or heteroaralkyl, arylsulfonyl, heteroarylsulfonyl, etc.; R1 = carboxyalkyl, alkoxycarbonylalkyl, N-(un)substituted carbamoylalkyl, qyanoalkyl, (un)substituted aralkyl or heteroaralkyl; R2 = (un)substituted lower alkyl; R3 = (un)substituted alkyl, alkeyl, alkynyl, cycloalkyl, cycloalkyl, or oxaaliph. (un)substituted or optionally oxidized cyclothioalkyl or cyclothioalkyl, R6 = H, (un)substituted lower alkyl; R5 = (un)substituted alkyl, alkoxy, cycloalkyl, or heterocyclyl, alkoxycarbonyl, cyano, (un)substituted arzbamoyl, (un)substituted aryl or heteroaryl, or CO2H where m is other than 0; R7 = H, alkoxy, (un)substituted cycloalkyl, cycloalkoxy, cycloalkenyloxy, aryl, heteroaryl, aryloxy, heteroaryloxy, aralkyloxy, heteroaryl, aryloxy, heteroaryloxy, aralkyloxy, heteroaryl, cycloalkenyloxy, aryl, heteroaryl, aryloxy, heteroaryloxy, aralkyloxy, heteroaryl, cycloalkyl, cycloalkyl,

L4 ANSWER 24 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 25 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) = H or Z'Z'' = O or S; Z1, Z2 = direct bond, O, S; Z3 = S02, direct bond; Z4 = direct bond, O, S, NH; Z5 = direct bond, (un) substituted lower alkenyl; m, n = 0, 1; p = 1-3; q = 0-5] or hydrate, solvate, N-oxide, or prodrug thereof or a pharmaceutically acceptable salt thereof are. They are esp. useful for inhibiting the prodn. or physiol. effects of tumor necrosis factor (TNF) and inhibit cAMP phosphodiesterase and are useful for the treatment of disease states assocd with abnormally high physiol. levels of cytokines such as TNF or those assocd with pathol. (e.g. ma

ma
as bronchodilators or inflammation) conditions that are modulated by
inhibiting enzymes such as cAMP phosphodiesterase (no data). In
particular, they are used for treating a disease state capable of being
modulated by inhibiting TNF, e.g., joint inflammation, arthritis,
rheumatoid arthritis, rheumatoid spondylitis and osteoarthritis, sepsis,
septic shock, gram neg. sepsis, toxic shock syndrome, acute respiratory
distress syndrome, asthma, bone resorption diseases, reperfusion injury,
graft vs. host reaction, allograft rejection malaria, myalgias, HIV,

cachexia, Crohn's disease, ulcerative colitis, pyresis, systemic lupus erythematosus, multiple sclerosis, type I diabetes mellitus, psoriasis, Behcet's disease, anaphylactoid purpura nephritis, chronic glomerulonephritis, inflammatory bowel disease, and leukemia. They are also used for treating a pathol. condition assocd. with a function of

phosphodiesterase, eosinophil accumulation or function of the eosinophil, phosphodiesterase, eosinophil accumulation or function of the eosinophil, e.g., asthma, atopic dermatitis, urticaria, allergic rhinitis, psoriasis, rheumatic arthritis, ulcerative colitis, Crohn's disease, adult respiratory distress syndrome, diabetes insipidus, keratosis, dermatitis, cerebral senility, multiinfarct dementia, senile dementia, memory impairment assocd. with Parkinson's disease, cardiac arrest, stroke, and intermittent claudication. The present invention is also directed to their pharmaceutical use, pharmaceutical compns. contg. the compds., and methods of their prepn. Thus, 2-(3-cyclopentyloxy-4-methoxyphenyl)-5-hydroxymethyl-2-(4-pyridylmethyl)indan-1,3-dione was treated with NaH in THF, tosylated by tosyl chloride at 0° to room temp. For 2 h, and then condensed with 1-methylpiperazine in the K2CO3 in acetone at room temp. for 4 days the presence of K2CO3 in acetone to give the title compd., piperazinylmethylpyridylmethylindandione deriv. (II). 201287-52-7P

20128/-52-79
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of substituted aromatic compds. as inhibitors of tumor

factor and cAMP phosphodiesterase) 201287-52-7 CAPLUS

Senzoic acid, 3-[(1-imino-2-phenylpropyl)amino]-4-methoxy-, methyl ester (CA INDEX NAME)

06/05/2008

10/568,760

Page 20

L4 ANSWER 25 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 26 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

PLUS COPYRIGHT 2008 ACS on STN
1998:31305 CAPLUS
128:102087
Substituted axabicyclic compounds and their use as
inhibitors of the production of TNF and cyclic AMP
phosphodiesterase
Cox, Paul Joseph; Bower, Shelley; Aldous, David John;
Astles, Peter Charles; McGarry, Daniel Gerard; Hulme,
Christopher; et al.
Regan, John Robinson, UK; Huang, Fu-Chih;
Rhone-Foulenc Rorer Ltd.; Cox, Paul Joseph; Bower,
Shelley; et al.
PCT Int. Appl., 355 pp.
CODEN: PIXXD2
Patent
English
1

INVENTOR(S):

PATENT ASSIGNEE(S):

SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

												LICAT						
												1997-						
		W:	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BF	, BY,	CA,	CH,	CN,	CU,	CZ,	DE,
			DK,	EE,	ES,	FI,	GB,	GE,	GH,	HU,	IL	, IS,	JP,	KE,	KG,	KP,	KR,	KZ,
												, MK,						
							SE,	SG,	SI,	SK,	SL	, TJ,	TM,	TR,	TT,	UA,	UG,	US,
				VN,														
		RW:										, BE,						
											SE	, BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,
								TD,										
												1997-						
												1997-						
	ZA	9705	446			A		1998	1221		ZA	1997- 1997-	5446			1	9970	619
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GP	, IT,	LI,	LU,	NL,	SE,	PT,	IE,
FI																		
	JP	2000	5097	19		T		2000	0802		JP	1998-	5025	03		J	9970	619
		6303				В1						1998-						
		6800						2004				2000-						
		2002						2002				2002-						
	US	2005	0038	069		AI		2005	0217		US	2004-	9330	7.7		-	0040	901
DD 7.0		7329				B2		2008	0212			1005						c 1 0
PRIO	RITY	APP:	LN.	INFO	. :						GB	1996-	1276	U		A 1	9960	619
											US	1996-	2304	7P		P 1	9960	802
											WO	1997-	GB16	39	,	W]	9970	619
											US	1998-	2163	92		A1 1	9981	218
											US	2000-	6125	30		A3 2	0000	707

OTHER SOURCE(S): MARPAT 128:102087

ANSWER 26 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

$$(\mathbb{R}^{1}\mathbb{Z}^{1})_{\underset{\mathbb{R}}{\mathbb{N}}} \times \mathbb{R}^{2}\mathbb{R}^{1} \times \mathbb{R}^{3} \qquad \mathbb{I} \qquad \mathbb{O}^{\mathbb{N}}$$

AB The invention is directed to physiol. active compds. of formula I [Wherein AB = fused bicyclic ring system, of approx. 10-13 ring members, wherein A = azaheterocycle ring and B = azaheterocryl or optionally halo-substituted benzene ring; R1 = H, (hydroxy- or halo-substituted) alkyl, and also alkenyl, alkynyl, or CHO when Z1 = bond; R2 = H, alkenyl, alkoxy, alkyl, aryl, aryloxy, cyano, etc.; R3 = wide variety of sidechains and functional groups; A1 = bond, (un) substituted alkylene, alkenylene, alkynylene; Z1 =

functional
groups; A1 = bond, (un) substituted alkylene, alkenylene, alkynylene; Z1 =
bond, O, S, NH; m, n = 0, 1; provided that (n+m) = 1] and their N-oxides,
prodrugs, and pharmaceutically acceptable salts and solvates. I inhibit
the production or physiol. effects of TNF, and inhibit cAMP
phosphodiesterase

(PDE IV). The invention is also directed to pharmaceutical compns. comprising I, their pharmaceutical use, and methods for their

comprising I, their pharmaceutuas use, and modeled in preparation For instance, 7-methoxy-2-(methoxymethyl)-3H-benzimidazole-4-carboxylic acid (preparation given) was treated with O-benzotriazol-1-yl-N,N',N' bis (tetramethylene) uronium tetrafluoroborate to give the 1-benzotriazolyl ester, which was amidated with 4-amino-3,5-dichloropyridine in THF (after treatment of the latter with Na diethylaluminate) to give the title

ound II. Compds. I had IC50 of 10-5 to 10-10 M against guinea pig macrophage PDE IV, with 50- to 10,000-fold selectivity for PDE IV vs. PDE I, II,

III.

L4 ANSWER 26 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

06/05/2008

ANSWER 27 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1996:464318 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 125:114673 125:21527a,21530a 125:21527a, 21530a
Preparation of benzyloxyphenylalkylbenzoates and
related compounds as analgesics and prostaglandin
antagonists
Breault, Gloria Ann; Oldfield, John; Tucker, Howard;
Warner, Peter
Zeneca Limited, UK
PCT Int. Appl., 172 pp.
CODEN: PIXXD2 TITLE: INVENTOR(S): PATENT ASSIGNEE(S): DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: A A A1 19960412 19960506 19960925 19991222 ZA 1995-8622 AU 1995-36162 EP 1995-933542 AU 9536162 EP 733033 EP 733033 19951012 19951012 В1 R: CH, DE, FR, GB, IT, LI
JP 09511529 T 19971118
US 5811459 A 19980922 JP 1995-513027 US 1996-647977 GB 1994-20557 19951012 US 5811459 PRIORITY APPLN. INFO.: A 19941012 W 19951012 WO 1995-GB2417 OTHER SOURCE(S): MARPAT 125:114673
AB Ortho-substituted Ph, naphthyl, and heterocyclic ethers (> 600 compds.) were prepared for use in treating pain mediated by the E-type prostaglandins prostagiandins (no data). Thus, 2-PhCH2OC6H4(CH2)3C6H4CO2H-4 was prepared from 2-HOC6H4Ac CSHARC
and 4-OCHCSH4CO2Me in 5 steps.
179256-35-0P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(preparation of benzyloxyphenylalkylbenzoates and related compds. as analgesics and prostaglandin antagonists)
179256-35-0 CAPLUS

TOTAGONETHALINGENINGE. M-methyl-2-(phenylmethoxy)- (CA INDEX NAME) analgesics and prostagrammin antagoniets,
179256-35-0 CAPLUS
Benzeneethanimidamide, α-methyl-2-(phenylmethoxy)- (CA INDEX NAME)

ACCESSION NUMBER: 1995;968831 CAPLUS

DOCUMENT NUMBER: 124:175546

ORIGINAL REFERENCE NO: 124:32547a,32550a

CITILE: Garigipati's reaction

AUTHOR(S): Garigipati's reaction

MOSS, Robert A.; Ma, Wei; Merrer, Dina C.; Xue, Song

CORPORATE SOURCE: Dep. Chem., Rutgers, The State Univ. New Jersey, New

Brunswick, NJ, 08903, USA

SOURCE: Tetrahedron Letters (1995), 36(48), 8761-4

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER: Elsevier

DOCUMENT TYPE: Journal

LANGUAGE: English

CTHER SOURCE(S): CASREACT 124:175546

AB Reaction with methylchloroaluminum amide readily converts sterically hindered nitriles, e.g., 1-adamantanecarbonitrile, to amidines.

I 173601-37-IP

RL: SFN (Synthetic preparation); PREP (Preparation) foreparation of amidines by Garigipati amination of sterically hindered nitriles)

RN 173601-37-I CAPLUS

CN Benzeneethanimidamide, α-methyl-α-phenyl- (CA INDEX NAME)

L4 ANSWER 27 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

ANSWER 29 OF 60 CAPLUS COPYRIGHT 2008 ACS ON STN

USSION NUMBER: 1995:638226 CAPLUS

MENT NUMBER: 123:95494

123:95494

123:95492h, 9983a

E: Preparation of propionic acid derivatives as serine protease inhibitors

NUTCR(S): Muramatsu, Mutsumi; Tamura, Toshiaki; Yanagi, Toshiharu

UNT ASSIGNEE(S): Teikoku Chemical Industries Co. Ltd., Japan

PCT Int. Appl., 98 pp.

COEN: PIXXD2

Patent

USAGE: Japanese

LY ACC. NUM. COUNT: 1

NUT INFORMATION: ACCESSION NUMBER:
DOCUMENT NUMBER:
ORIGINAL REFERENCE NO.: INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. KIND DATE W: JP, KR, US
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
EP 673924
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, PT, SE
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, A 19921210 PRIORITY APPLN. INFO.: JP 1992-360711

> JP 1993-318909 A 19931112 WO 1993-JP1783 W 19931209

OTHER SOURCE(S): MARPAT 123:55494

2-[P-(p-guanidinobenzoyloxy)phenyl]propionic acid derivs. represented by general formula [I, A = OH, Cl-6 lower alkoxy, NRIR2, Cl-8 lower alkoxy which may be substituted by halogen, optionally substituted aryl, COB or succinimido; Rl, R2 = H, Cl-8 lower alkyl, optionally substituted aralkyl or alternatively Rl and R2 are combined together with the adjacent nitrogen atoms to represent a heterocycle; B = OH, Cl-8 lower alkyl, optionally substituted aryl, optionally substituted aryloxy, Cl-8 lower alkoxy, optionally substituted aralkyloxy, NRIR2 (wherein Rl and R2 are each as defined above)] or pharmaceutically acceptable acid-addition s

salts
thereof is prepared These compds. are useful as inhibitors of serine
protease such as trypsin, chymotrypsin, plasmin, or thrombin and for the
treatment of pancreatitis, bleeding, thrombosis, nephritis, and general
internal clot and prevention of blood coagulation under perfusion during
dialysis or exchange of blood plasma. Thus, 3.44 g DCC was added to a
mixture of 3.85 g N,N-dimethylcarbamoylmethyl
2-(4-hydroxyphenyl)propionate,

06/05/2008

10/568,760

14 ANSWER 29 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)
3.00 g 4-guanidinobenzoic acid hydrochloride, and 20 ml pyridine and the resulting mixt. was stirred at room temp. overnight to give, after workup and acidification with MeSO3H, N,N-dimethylcarbamoylmethyl
2-[4-(4-guanidinobenzoyloxy)phenyl]propionate methanesulfonate, which in vitro showed IC50 of 1.4 + 10-7 and 1.9 + 10-8 M against trypsin and plasmin, resp. A tablet formulation contg. (S)-(+)-I.MeSO3H (A = CH2Ph) was prepd.

IT 159239-63-IP
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study unclassified): SDN (Superhetic account)

Page 22

logical study, unclassified); SFN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of [(guanidinobenzoyloxy)phenyl]propionic acid derivs. as serine protease inhibitors) 159239-63-1 CAPLUS Benzoic acid, 4-[(aminoiminomethyl)amino]-, 4-(2-amino-2-imino-1-methylethyl)phenyl ester, dimethanesulfonate (9CI) (CA INDEX NAME)

CRN 159239-62-0 CMF C17 H19 N5 O2

CM

CRN 75-75-2 CMF C H4 03 S

L4 ANSWER 30 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

160154-97-2 CAPLUS
Benzeneethanimidic acid, α-hydroxy-α-methyl-,
cyclopentylidenehydrazide (9CI) (CA INDEX NAME)

160154-98-3 CAPLUS Benzeneethanimidic acid, $\alpha\text{-hydroxy-}\alpha\text{-methyl-},$ (phenylmethylene)hydrazide (9CI) (CA INDEX NAME)

ANSWER 30 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1995:224478 CAPLUS

ACCESSION NUMBER:

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 122:81192 122:15427a,15430a

122:15427a,15430a
4-hydrazonooxazolidin-2-ones from @-substituted
glycolamidrazones
Geffken, D.; Holst, C.
Inst. Pharmazie, Universitaet Hamburg, Germany
Pharmazie (1994), 49(11), 821-4
CODEN: PHARAT; 1SSN: 0031-7144
Govi-Verlag Pharmazeutischer Verlag
Journal TITLE:

CORPORATE SOURCE: SOURCE:

Journal

AUTHOR (S):

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
GI German CASREACT 122:81192

Hydrazinolysis of the glycolimidates gave glycolamidrazones which were with acetone or benzaldehyde to give hydrazono derivs. of type 4. I (R1 = alkyl, Ph, etc.; R2 = H, Me). R3 = Me, Ph, etc.; R4 = H, Me). Cyclic carbonylation of I with 1,1'-carbonylatimidazole yields 4-hydrazono-2-oxazolidinones II (same R1-R4). 160154-90-5P, a-Hydrazyow-a-methylbenzeneethanimidic acid hydrazide 160154-94-9P 160154-97-2P 160154-98-3P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant) research

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); (Reactant or reagent) (Reactant or reagent) (Synthetic preparation of (hydrazono)xazolidinones from glycolamidrazones) 160154-90-5 (APLUS Benzeneethanimidic acid, α-hydroxy-α-methyl-, hydrazide (CA INDEX NAME)

160154-94-9 CAPLUS Benzeneethanimidic acid, α -hydroxy- α -methyl-, (1-methylethylidene)hydrazide (9CI) (CA INDEX NAME)

L4 ANSWER 31 OF 60 CA
ACCESSION NUMBER:
DOCUMENT NUMBER:
ORIGINAL REFERENCE NO.:
TITLE:
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:

ANSWER 31 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
SSION NUMBER: 1994:692802 CAPLUS
MENT NUMBER: 121:292802
121:53304h, 53305a,53307a
Amidinoethyl derivative
NTOR(S): Amidinoethyl derivative
NTOR(S): Teikoku Hormone Mfg Co Ltd, Japan
Jpn. Kokai Tokkyo Koho, 4 pp.
CCE: JRY AKXAF
MENT TYPE: CODEN: JKXXAF
MENT TYPE: Japanese
LY ACC. NUM. COUNT: 1
Japanese
LY ACC. NUM. COUNT: 1

DOCUMENT TYPE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. JP 06228078 PRIORITY APPLN. INFO.: JP 1993-305706 JP 1993-305706 19931028 19931028 19940816

$${\rm H_{2}NC\;(NH)\;CHMe} \xrightarrow{\hspace*{1cm}} {\rm OCO} \xrightarrow{\hspace*{1cm}} {\rm NHC\;(NH)\;NH_{2}}$$

Amidinoethyl derivative I or its salts are useful as serine protease inhibitors for treatment of diseases (e.g. inflammation, cardiovascular diseases, and pancreatic diseases), caused by abnormalities of the

enzyme.
4-(1-Amidinoethyl)phenol methanesulfonic acid salt (preparation given)

II inhibited trypsin and thrombin with IC50 of 3.2 + 10-7 and 6.3

+ 10-9 (no unit given). 159239-62-0P 159239-63-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological $\,$

serine protease)
159239-62-0 CAPLUS
Benzoic acid, 4-[(aminoiminomethyl)amino]-, 4-(2-amino-2-imino-1-methyl)phenyl ester (CA INDEX NAME)

06/05/2008 Habte

L4 ANSWER 31 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

159239-63-1 CAPLUS
Benzoic acid, 4-[(aminoiminomethyl)amino]-, 4-(2-amino-2-imino-1-methylethyl)phenyl ester, dimethanesulfonate (9CI) (CA INDEX NAME)

CRN 159239-62-0 CMF C17 H19 N5 O2

CM

CRN 75-75-2 CMF C H4 O3 S

L4 ANSWER 32 O OTHER SOURCE(S): ANSWER 32 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN R SOURCE(S): MARPAT 121:102030 (Continued)

AB The N-arylhydrazine derivs. I [A,B,W=N,CR4;Y=halo,CN,NO2,(halo)alkyl,(halo

[A, B, W=M, CR4)Y=halo, CM, NO.2, (halo)alkyl, (halo)
 lakoxy,n=0,1,2;Q=NR2CRO,N:CRX1,N:CR (NR3R4);R=H, (halo)alkyl,cycloalkyl, (halo)alkoxy,etc.;RI, R2=H, alkyl;R3,R4=H, (un) substituted alkyl;Ph or pyridyl, etc.] are prepared as acaricides and insecticides. Treatment of 2,6-dichloro-4(rrifluoromethyl) phenylhydrazine with trimethylacetyl chloride, in Cl2CH2, gave 2,2-dimethylpropionic acid 2-(2,6-dichloro-α,α,α-trifluoro-p-tolyl) hydrazide (II). Lima bean leaves dipped in 300 ppm II were lethal to Southern armyworm (Spodoptera eridania) 3rd instar larvee.
 IT 156820-05-2P 156820-21-2P 156820-27-8P
 RLi AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSI (Biological study, unclassified); SPN (Synthetic preparation); BIO. (Biological study); PREF (Preparation); USES (Uses) (preparation of, as acaricide and insecticide)
 RN 156820-05-2 CAPLUS
 CN Benreneethanimidic acid, N-ethyl-α,α-dimethyl-, 2-(2,6-dichloro-4-(trifluoromethyl)phenyl]hydraride (CA INDEX NAME)

 $156820-21-2 \quad CAPLUS \\ Benzeneethanimidic acid, \\ \alpha,\alpha-dimethyl-N-(2-phenylethyl)-, \\ 2-(2,6-dih)ozc-4-(trifluoromethyl)phenyl]hydrazide \quad (CA INDEX NAME)$

ANSWER 32 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1994:502030 CAPLUS

ACCESSION NUMBER:

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 121:102030 121:18219a,18222a

TITLE:

Nearylhydrazine derivatives as insecticides and acaricides. Furch, Joseph Augustus; Kuhn, David George; Hunt, David Allen; Lew, Albert Chieh; Gronostajski, Cynthia INVENTOR(S):

David Allen; Lew, Albert Ch Emma American Cyanamid Co., USA Eur. Pat. Appl., 50 pp. CODEN: EPXXDW Patent English 1 PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT INFORMATION:				
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 604798	A1	19940706	EP 1993-119754	19931208
EP 604/98	BT DA	20020220	GB, GR, IE, IT, LI,	THE NUMBER OF
US 342U163	A.	19930330	US 1992-998105 AT 1993-119754	19921229
MI 213307	1 7	20020315	ES 1993-119754	19931200
ES 21/3088	T3	20021016	CZ 1993-119754 CZ 1993-2808	19931208
CZ 2004/9	30	200000412	CZ 1993-2000	19931217
MU 9352679	M.	19940714	AU 1993-32679	19931224
AU 6/5255	B2	19970130	AU 1993-52679 CA 1993-2112420 RO 1993-1796	10071004
CA 2112420	C	20070213	CA 1993-2112420 RO 1993-1796 SK 1993-1484	19931224
KO 113556 SK 281733	BI	19980828	KO 1993-1/96	19931227
SA 201733	30	20010710	IL 1993-108188	19931227
			CN 1993-100100	
CN 1089938 CN 1044600			CN 1993-121610	19931228
CN 1044600	ь	19990811		
ZA 9309740	A	19940818	ZA 1993-9740	19931228
JP 06293605	A	19941021	JP 1993-350030 BR 1993-5254 HU 1993-3772	19931228
JP 3816543	B2	20060830	PP 1003 F0F4	10071000
BK 93U5254	A	19941101	BK 1993-5254	19931228
HU 6/294 HU 221126	AZ	19950328	HU 1993-3772	19931228
			PL 1993-317481	
			PL 1993-301659	
RU 2140738	C1	19991110	RU 1993-56849	19931228
CA 211242U	AI	19940630	CA 1994-2112420	19940121
US 5585389	A	19961217	CA 1994-2112420 US 1995-431227 US 1995-431154 US 1995-430631 JP 2005-134574	19950428
US 5646278	A	19970708	US 1995-431154	19950428
US 5693860	A	19971202	US 1995-430631	19950428
JP 2005263809	A	20050929	JP 2005-134574	20050502
PRIORITY APPLN. INFO.:			US 1992-998101	A 19921229
			US 1992-998104	A 19921229
			US 1992-998105	* 10001000
			09 1337-338103	W 12351553
			JP 1993-350030	A3 19931228

ANSWER 32 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) 156820-27-8 CAPLUS Benzeneethanimidic acid, 4-chloro-N-ethyl- α , α -dimethyl-, 2-[2,6-dichloro-4-(trifluoromethyl)phenyl]hydrazide (CA INDEX NAME)

06/05/2008 Habte

ANSWER 33 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1991:6034 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 114:6034 114:1187a,1190a

Preparation of N-hydroxyamidines as acaricides and agricultural and horticultural fungicides Kishimoto, Takashi; Hayakawa, Koichi; Nakayama, TITLE: INVENTOR(S):

Yamada, Tomio; Takahashi, Eiko; Hashimoto, Akira; Sano, Shinsuke; Hosokawa, Hiroyasu Nippon Soda Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 29 pp. CODEN: JKXXAF Patent Japanese 1 PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 02006453 PRIORITY APPLN. INFO.:	A	19900110	JP 1988-158393 JP 1988-158393	19880627 19880627

OTHER SOURCE(S): MARPAT 114:6034

мосн2с≡сн

Amidines R1[R5(R2O)N]C:NR3 (I) and R1(R3R4N)C:NOR2 II [R1 = H, (un)substituted Ph, alkyl optionally substituted by (un)substituted Ph, naphthyl, alkylthio, aralkylthio, (un)substituted NH2, cyclic amino, (un)substituted heterocyclyl, R2 = H, (un)substituted alkyl, alkenyl, alkynyl, XR6, X = CO, CONH, COZ, COCO, R6 = alkyl, (un)substituted alkyl, alkenyl, Ph, or aralkyl, P(:Y)(CR7)2, Y = O, S, R7 = alkyl, R3 = H, (un)substituted alkyl, alkynyl, ZR8; Z = CO, CS, COZ, COCO, CONH, SOZ, OZC; R8 = (un)substituted alkyl, alkenyl, or aralkyl, piperidino; R4 = H, alkyl; R5 = alkyl, aralkyl, (un)substituted aralkylcarbonyl] are pared,

Hea, by reaction of R1C(X):NOR2 (X = halo) with HNR3R4. Thus, PhCH2CCC1 was added to a solution of 2,6-C12C6H3C(NH2):NOCH2C.tplbond.CH in

ene and the mixture was refluxed overnight to give a benzamidine III. A total of 574 II were prepared and 18 II at 125 ppm completely controlled

5/4 II wese Parparon Tetranychus

Tetranychus

urticae and III and 46 others at 200 ppm controlled 77-100% Erysiphe

gramınıs. 129860-61-3P 129860-62-4P 129860-63-5P

ANSWER 33 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

129860-67-9 CAPLUS

Benzeneacetamide, 4-methoxy-N-[2-phenyl-1-[(2-propynyloxy)amino]propylidene]- (9CI) (CA INDEX NAME)

129860-68-0 CAPLUS
Benzeneacetamide, N-[1-(ethoxyamino)-2-phenylpropylidene]-4-methoxy-

ANSWER 33 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) 129860-64-6F 129860-67-9P 129860-68-0P RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SFN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of, as acaricide and agrochem. fungicide) 129860-61-3 CAPLUS Benzeneacetamide, 4-methoxy-N-[2-methyl-2-phenyl-1-[(2-propynyloxy)amino]propylidene]- (9CI) (CA INDEX NAME)

129860-62-4 CAPLUS
Benzeneacetamide, N-[1-(ethoxyamino)-2-methyl-2-phenylpropylidene]-4methoxy- (981) (CA INDEX NAME)

RN

129860-63-5 CAPLUS
Benzeneacetamide, 4-(1-methylethoxy)-N-[2-methyl-2-phenyl-1-[(2-propynyloxy)amino]propylidene]- (9CI) (CA INDEX NAME)

129860-64-6 CAPLUS Benzeneacetamide, 4-methoxy-N-[2-(4-methoxyphenyl)-2-methyl-1-[(2-propynyloxy)amino]propylidene]- (9CI) (CA INDEX NAME)

L4 ANSWER 34 OF 60 C.
ACCESSION NUMBER:
DOCUMENT NUMBER:
ORIGINAL REFERENCE NO.:

ANSWER 34 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

ISSION NUMBER: 1990:77194 CAPLUS
INDERT NUMBER: 112:77194
INDERT NUMBER: 112:17194
INDERT 112:13203a, 13206a
Freparation of oxadiazoles as central muscarinic acetylcholine receptor stimulants and pharmaceutical compositions containing them
ENTOR(S): Baker, Raymond; Merchant, Kevin J.; Saunders, John;
Street, Leslie J.

Merck Sharp and Dohme Ltd., UK
Eur. Pat. Appl., 27 pp.
CODEN: EPXXDW
Patent
English
ILIT ACC. NUM. COUNT: 1

INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA'	TENT	NO.			KINI	D.F	ATE		API	PLICATION :	NO.		DATE
												-	
EP	3238	64			A2	19	890712		EP	1989-2000	01		19890102
EP	3238	64			A3	19	911218						
	R:	ΑT,	BE,	CH,	DE,	ES, E	R, GB,	GR,	, I.	r, LI, LU,	NL, SE		
ZA	8900	045			A	19	900926		ZA	1989-45			19890104
DK	8900	041			A	19	890709		DK	1989-41			19890106
AU	8927	798			A	19	890720		AU	1989-2779	8		19890106
AU	6283	11			B2	19	920917						
JP	0214	9580			A	19	900608		JP	1989-571			19890106
PRIORIT	Y APP	LN.	INFO	. :					GB	1988-394		A	19880108
									GB	1988-1351	3	Α	19880608
									GB	1988-2489	8	A	19881024

AB The title compds. [I; R1 = non-aromatic aza(bi)cyclic ring residue, e.g., pyrrolidinyl, piperidinyl, tetrahydropyridinyl; R2 = (substituted) saturated hydrocarbyl, e.g., Pr, Me2CH; one of X, Y, and Z = O and the other 2 =

central muscarinic acetylcholine receptor stimulants, useful for

treatment and prevention of neurodegenerative diseases, are prepared via cyclocondensation of R3CO2H with HON:CR4NH2 or R4CONHNH2 [one of R3 and = non-aromatic aza(bi)cyclic ring residue and the other = (substituted)

saturated
hydrocarbyl]. PhCH2C(NH2):NOH was condensed with 3(methoxycarbonyl) quinuclidine in THF containing NaH to give
3-(3-benzyl-1,2,4-oxadiazol-5-yl)quinuclidine, isolated as its

L4 ANSMER 34 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) hemioxalate. A tablet comprising
3-(3-eyelopropyl-1,2,4-oxadiazol-5-yl)-1- arabicylo(2,2.1)heptane 1.0, microcryst. cellulose 49.25, modified food corn starch 49.25, and Mg stearate 0.50 mg was formulated. I had an IC50 of better than 10 μM for displacement of specifically bound [3H]-N-methylscopolamine from muscarinic receptors of rat cortical membrane prepns.

IT 42191-51-5, 2-Phenylpropionamide oxime
RL: RCT (Reactant); RRCT (Reactant or reagent) (reaction of, in preparation of oxadiazoles for treatment of neurodegenerative diseases)

RN 42191-51-5 CAPLUS
CN Benzenethanimidamide, N-hydroxy-α-methyl- (CA INDEX NAME)

ANSWER 35 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN 55769-91-0 CAPLUS Benzeneethanimidamide, N-(aminocarbonyl)- α -methyl-, monohydrochloride (9CT) (CA INDEX NAME) (Continued)

78622-01-2 CAPLUS Benzeneethanimidamide, N-(aminocarbonyl)-2,6-dichloro- α -methyl-, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

55770-09-7P 78622-11-4P 78630-47-4P

55/70-U9-19 / Pos22-II-4P / Ro3U-4/-4P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation and hydration of, N-carbamoylamidine analog from)
55770-09-7 CAPLUS
Benzeneethanimidamide, 2-chloro-N-cyano-α-methyl- (CA INDEX NAME)

78622-11-4 CAPLUS Benzeneethanimidamide, N-cyano- α -methyl- (CA INDEX NAME)

78630-47-4 CAPLUS Benzeneethanimidamide, 2,6-dichloro-N-cyano-α-methyl- (CA INDEX

ANSWER 35 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1981:480429 CAPLUS

ACCESSION NUMBER:

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.:

TITLE:

1981:480429 CAPLUS
95:80429
95:13591a,13594a
Synthesis and properties of the tremor-inducing
N-carlamoylacetamidine derivative LON-954 and some
related compounds
Bream, John B.; Picard, Claude W.; White, Trevor G.
Wander Res. Inst., Wander Ltd., Bern, CH-3001, Switz.
European Journal of Medicinal Chemistry (1981), CORPORATE SOURCE: SOURCE: 16(2).

173-9 CODEN: EJMCA5; ISSN: 0009-4374 Journal English CASREACT 95:80429

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

$$zc = NCONR^{1}R^{2})NR^{3}R^{4}$$

The hydration of N-cyanophenylacetamidines gave N-carbamoyl analogs I (Z

ANSWER 35 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

78622-20-5P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and reaction of, with cyanogen bromide) 78622-20-5 CAPLUS Benzeneethanimidamide, 2,6-dichloro-α-methyl- (CA INDEX NAME)

78622-19-2 RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of cyanogen bromide with free base from) 78622-19-2 CAPLUS Benzeneethanimidamide, 2,6-dichloro- α -methyl-, monohydrochloride (9C1) (CA INDEX NAME)

• HCl

55770-08-6 78622-24-9 RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with oyanogen bromide) 55770-08-6 CAPLUS Benzeneethanimidamide, 2-chloro- α -methyl-, monohydrochloride (9CI) (CA INDEX NAME)

L4 ANSWER 35 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

• HCl

78622-24-9 CAPLUS Benzeneethanimidamide, α -methyl-, monohydrochloride (9CI) (CA INDEX NAME)

• HCl

L4 ANSWER 36 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 1978:406123 CAPLUS
DOCUMENT NUMBER: 89:6123
ORIGINAL REFFERENCE NO: 89:1043a, 1046a
TITLE: 1NVENTOR(S): 7akacs, Kalman; Nagy, Peter Literati; Kiss, Ilona; Sinay, Antal; Szentivanyi, Matyas; Virag, Sandor; Farago, Katalin
PATENT ASSIGNEE(S): Chionin Gyogyszer es Vegyeszeti Termekek Gyara Rt., Hung.
SOURCE: GET. Offen., 33 pp.
CODEN: GWXXEX
DOCUMENT TYPE: Fatent German
FATENT INFORMATION: TO THE CONTROL OF THE CONT

PATENT NO.	KIND	DATE	AP	PLICATION NO.	DATE
DE 2738589	A1	19780302	DE	1977-2738589	1977082
DE 2738589	C2	19900419			
HU 19948	A2	19810528	HU	1976-CI1682	1976082
HU 177578	В	19811128			
AT 7706054	A	19790815	AT	1977-6054	1977082
AT 355554	В	19800310			
SE 7709482	A	19780228	SE	1977-9482	1977082
SE 435280	В	19840917			
SE 435280	C	19841220			
NL 7709276	A	19780301	NL	1977-9276	1977082
NL 187478	В	19910516			
NL 187478	C	19911016			
IL 52804	A	19810629	IL	1977-52804	1977082
DD 132433	A5	19780927	DD	1977-200719	1977082
CS 204008	B2	19810331	CS	1977-5551	1977082
GB 1582029	A	19801231	GB	1977-35745	1977082
BE 858134	A1	19771216	BE	1977-180447	1977082
DK 7703797	A	19780228	DK	1977-3797	1977082
DK 150196	В	19870105			
DK 150196	C	19870706			
FI 7702551	A	19780228	FI	1977-2551	1977082
FI 68396	В	19850531			
FI 68396	C	19850910			
NO 7702958	A	19780228	NO	1977-2958	1977082
NO 144793	В	19810803			
NO 144793	C	19811111			
FR 2362845	A1	19780324	FR	1977-26070	1977082
FR 2362845	B1	19810109			
JP 53050131	A	19780508	JP	1977-102504	1977082
JP 62016942	В	19870415			
AU 7728254	A	19790301	AU	1977-28254	1977082
AU 521432	B2	19820401			
PL 106317	B1	19791231	PL	1977-200480	1977082
PL 107628	B1	19800229	PL	1977-206476	1977082
SU 730296	A3	19800425	SU	1977-2514754	1977082
CA 1077506	A1	19800513	CA	1977-285529	1977082

L4	ANSWER 36 OF (50 CAPLUS	COPYRIGHT 200	8 ACS on STN	(Continued)
	CH 630344	A5	19820615	CH 1977-10473	19770826
	US 4187220	A	19800205	US 1977-829148	19770830
	CS 204009	B2	19810331	CS 1978-5952	19780914
	AT 7808741	A	19800815	AT 1978-8741	19781207
	AT 361457	В	19810310		
	US 4308399	A	19811229	US 1979-54791	19790705
PRIO	RITY APPLN. IN	FO.:		HU 1976-CI1682	A 19760827
				HU 1977-CI1682	A 19770426
				AT 1977-6054	A 19770822
				CS 1977-5551	19770824
				US 1977-829148	A3 19770830

OTHER SOURCE(S): MARPAT 89:6123

AB RRINCH2CH(OB)CH2ON:C(NH2)(CHR2)nr(CHR3)mR4 I (R = H, C1-5 alkyl; R1 = C1-5 alkyl, cycloalkyl, Ph, optionally substituted by OH or Ph; RRIN = heterocycle; R2 = H, C1-4 alkyl, Ph; R3 = H, C1-4 alkyl, cycloalkyl or

optionally substituted by halogen; R4 = optionally substituted

optionally substitutes of, cycloalkyl, aromatic or heterocyclic group; m = n = 0, 1, 2) and their salts were

ared Thus, PhC(NH2):NOH reacted with 1-chloro-3-piperidino-2-propanol in EtOH to give I (RRIN = piperidino, R4 = Ph, m = n = 0). I are useful as antidiabetics. 6661-55-0p

66611-55-0P
RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)
66611-55-0 CAPLUS
Benzeneethanimidamide, N-[2-hydroxy-3-[(1-methylethyl)amino]propoxy]α-methyl-, dihydrochloride (9CI) (CA INDEX NAME)

●2 HCl

(Continued)

L4 ANSWER 36 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

ANSWER 37 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1976:508660 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 85:108660 85:17445a,17448a 85:17445a,17448a Pyrimidine derivatives Komori, Saburo Yanagida, Shozo, Japan Jpn. Tokkyo Koho, 3 pp CODEN: JAXXAD Patent TITLE: INVENTOR(S): PATENT ASSIGNEE(S):

DOCUMENT TYPE. LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE JP 50037671 PRIORITY APPLN. INFO.: 19751204 JP 1970-35552 JP 1970-35552 19700425

For diagram(s), see printed CA Issue. Amidines I (R, Rl = Cl, alkyl, aralkyl, aryl) were heated with COC12 to give pyrimidines II (R2 = Cl, OH). Thus, 2.3 g I (R = Cl, Rl = Me), 2.3

CCC12, and PhCl were heated 90 hr in a sealed tube at 100-110 $^{\circ}$ to give 2.01 g II (R = R2 = C1, R1 = Me). Similarly prepared were II (R,

R2 given): C1, C1, OH; C1, Ph, C1; Me, Ph, C1; Et, Et, C1. 40645-76-9

406.45-76-9
RL: RCT (Reactant); RACT (Reactant or reagent) (cyclization of, with phosgene, pyrimidine derivative from) 406.45-76-9 CAPLUS
Benzeneethanimidantide, N-(1-chloro-2-phenyl-1-propen-1-yl)-α-methyl-, hydrochloride (1:1) (CA INDEX NAME)

HCl

ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: TITLE:

ANSWER 39 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

SSION NUMBER: 1976:30902 CAPLUS

MENT NUMBER: 84:5045a,5048a

Substituted α-phenylcarboxylic acids and their functional acid derivatives

E: nunctional acid derivatives

Rossi, Alberto

Ciba-Geigy A.-G., Switz.

CE: Patentachrift (Switz.), 6 pp. Division of Swiss

559,173.

CODEN: SWXXAS

MENT TYPE: Patentachrift

SUAGE: German

LLY ACC. NUM. COUNT: 1

INVENTOR(S).

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CH 566311	A5	19750915	CH 1971-13160	19690605
PRIORITY APPLN. INFO.:			CH 1971-13160	19690605

piperidinone

ridinone
I. Antiinflammatory I was effective on rat paws in the Kaolin edema test in oral doses of 30-100 mg/kg.
41789-12-2P
RL: SFN (Synthetic preparation); PREP (Preparation)

(preparation of) 41789-12-2 CAPJUS Benzeneethanimidamide, N-hydroxy-α-methyl-4-(1-methyl-6-oxo-3-piperidinyl)- (CA INDEX NAME)

ANSWER 38 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1976:508659 CAPLUS

ACCESSION NUMBER:

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 85:108659 85:17445a,17448a

TITLE: INVENTOR(S):

85:1/445a,1/448a Barbituric acid derivatives Komori, Saburo Yanagida, Shozo, Japan Jpn. Tokkyo Koho, 3 pp. CODEN: JAXXAD PATENT ASSIGNEE(S):

DOCUMENT TYPE. DOCUMENT TYPE: Patent
LANGUAGE: Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE В JP 50037673 PRIORITY APPLN. INFO.: 19751204 JP 1970-80192 JP 1970-80192

For diagram(s), see printed CA Issue. Nitriles RCHRICN(I) [R, Rl = (substituted) alkyl, Ph] or amides RCHRICNH2 (II) were treated with CCCl2 in the presence of HCl followed

treating the product with H2O to give III, which were also prepared by treating amidines IV (a mixture of cis and trans isomers) [R2, R3 = (substituted) alkyl, Ph] with CCC12 and then with H2O. Thus, 1.5 g

(R - R3 = Me) and 2.4 g COC12 in PhC1 were heated in a sealed tube 20 hr at 100-110° to give 0.23 g III (R = R1 = Me), which was also prepared by heating a mixture of isobutyronitrile, HC1, COC12 and PhC1 in a sealed tube 77 hr at 100-110°. Similarly prepared were III (R, R1 given): Me, Et; Me, Ph; Bu, CH2CH2Br; Et, m-OZNC6H4.
40645-76-9
RL: RCT (Reactant); RACT (Reactant or reagent) (cyclization of, with phosgene) 40645-76-9 CAPLUS
Benzeneethanimidamide, N-(1-chloro-2-phenyl-1-propen-1-yl)-\alpha-methyl-, hydrochloride (1:1) (CA INDEX NAME)

● HC1

ANSWER 40 OF 60 CAPLUS COPYRIGHT 2008 ACS ON STN
SSION NUMBER: 1975:155896 CAPLUS
MENT NUMBER: 82:155896
E: 82:24865a, 24868a
Araliphatic acetamidines
NTOR(S): Bream, John B.
NT ASSIGNEE(S): Dr. A. Wander, A.-G., Switz.
CE: Ger. Offen., 28 pp.
CODEN: GWXEN
MENT TYPE: German
LY ACC. NUM. COUNT: 1
NT INFORMATION:

L4 ANSWER 40 OF 60 CA
ACCESSION NUMBER:
DOCUMENT NUMBER:
ORIGINAL REFERENCE NO.:
TITLE:
INVENTOR(S):
SOURCE:
SOURCE:

DOCUMENT TYPE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2439299	A1	19750306	DE 1974-2439299	19740816
FR 2241300	A1	19750321	FR 1974-27773	19740809
FI 7402392	A	19750221	FI 1974-2392	19740812
NO 7402887	A	19750221	NO 1974-2887	19740812
SE 7410281	A	19750221	SE 1974-10281	19740812
DK 7404280	A	19750428	DK 1974-4280	19740812
NL 7410987	A	19750224	NL 1974-10987	19740816
DD 116606	A5	19751205	DD 1974-180552	19740816
BE 818988	A1	19750219	BE 1974-147735	19740819
JP 50052043	A	19750509	JP 1974-94329	19740819
AU 7472496	A	19760219	AU 1974-72496	19740819
ZA 7405340	A	19760331	ZA 1974-5340	19740820
PRIORITY APPLN. INFO.:			GB 1973-39263	A 19730820

GB 1973-44372 A 19730921

Thirty-three RnC6H5-nXC(:NR1)NHCONR2R3 (Rn = e.g. 3,4-Cl2, 3,4-Me2, 2-Cl, or 3-CF3; X = CH2, CHMe, or CH2CH2; R1-R3 = H or Me), useful as antidepressants, were prepared by hydrolysis of RnC6H5-nXC(:NR1)NHCN or

reaction of RnC6H5-nXC(:NR1)NH2 with R2NCO (R2 = e.g. Me) or with

IT

RN

reaction of RnC6Hb-nXC(:NRI)NH2 with R2NCO (R2 = e.g. Me) or with RZR3NCOCI.
55770-08-6
RL: RCT (Reactant); RACT (Reactant or reagent) (cyanation of)
55770-08-6 CAPLUS
Benzeneethanimidamide, 2-chloro-α-methyl-, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

06/05/2008

ANSWER 40 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) 55770-09-7P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and hydrolysis of) 55770-09-7 CAPLUS Benzeneethanimidamide, 2-chloro-N-cyano-\alpha-methyl- (CA INDEX NAME)

55769-76-1P 55769-81-8P 55769-91-0P 55769-95-4P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, for antidepressants) 55769-76-1 CAPLUS Benzeneethanimidamide, N-(aminocarbony1)-2-chloro-α-methyl-, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

55769-81-8 CAPLUS Benzeneethanimidamide, N-(aminocarbony1)-3,4-dichloro- α -methyl-, monbydrochloride (9C1) (CA INDEX NAME)

L4 ANSWER 40 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

● HCl

55769-91-0 CAPLUS Benzeneethanimidamide, N-(aminocarbonyl)- α -methyl-, monohydrochloride (9CI) (CA INDEX NAME)

• HCl

55769-95-4 CAPLUS Benzeneethanimidamide, N-(aminocarbonyl)-3,4-dimethoxy- α -methyl-, monohydrochloride (9CI) (CA INDEX NAME)

#C1

ANSWER 41 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
SSION NUMBER: 1973:466039 CAPLUS
MENT NUMBER: 79:10667a,10670a
E. Aromatic acetamindoxime O-carbamates
NTOR(S): Henderson, Rosetta M.
NT ASSIGNEE(S): du Pont de Memours, E. I., and Co.
U.S., 8 pp.
CDE: U.S., 8 pp.
TABLE TYPE: Patent
UNGE: Patent
English
LY ACC. NUM. COUNT: 1 L4 ANSWER 41 OF 60 CA ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: TITLE: INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. US 3742056 PRIORITY APPLN. INFO.: A US 1971-135806 US 1971-135806 19730626

Antihypertensive and antiinflammatory acetamidoxime O-carbamates, RnC6H5-nCHR1C(NH2):NO2CNHR2 (Rn = H, 4-Cl, 4-F, 2-Me, 4-MO2, 3,4-(MeO)2, 3,4-Me2, 2,4,6-Me3; R1 = H, Me; R2 = Me, Pr) were prepared by treating

acetamidoximes RnC6H5-nCHR1C(NH2):NOH with the isocyanates R2NCO.

42191-44-6P RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)
42191-44-6 CAPLUS
Benzeneethanimidamide, α-methyl-N-[[(methylamino)carbonyl]oxy]-,
monohydrochloride (9CI) (CA INDEX NAME)

● HCl

42191-51-5

%2191-31-3
%L: RCT (Reactant); RACT (Reactant or reagent)
 (reaction of, with alkyl isocyanates)
42191-51-5 CAPLUS
Benzeneethanimidamide, N-hydroxy-α-methyl- (CA INDEX NAME)

ANSWER 42 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

SSION NUMBER: 1973:418585 CAPLUS

MENT NUMBER: 79:18585

Substituted \(\alpha \) - phenylcarboxylicacids

NITOR (S): Rossi, Alberto

Ciba-Geigy A.-G.

Patentschrift (Switz.), 7 pp.

CODEN: SWXXAS

Patent

SUAGE: German

LLY ACC. NUM. COUNT: 1

NUM T INFORMATION: L4 ANSWER 42 OF 60 CA
ACCESSION NUMBER:
DOCUMENT NUMBER:
ORIGINAL REFERENCE NO.:
TITLE:
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND APPLICATION NO. A CH 534680 PRIORITY APPLN. INFO.: CH 1972-3553 CH 1972-3553 19730430

For diagram(s), see printed CA Issue. The piperidinylphenylpropionic acids I [R = 1-acetyl-2(or 4)-piperidinyl, 1-methyl-2-oxo-4(or 5, or 6)-piperidinyl; R1 = H, Et] were prepared Thus 4-(4-piperidinyl)phenylacetic acid was acetylated and then methylated

Buli-MeI to I (R = 1-acetyl-4-piperidinyl, RI = H). I were antiinflammatory at 30-100 mg/kg orally in the rat paw edema test. 41789-12-2P
RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 41789-12-2 CAPJUS
Benzeneethanimidamide, N-hydroxy-α-methyl-4-(1-methyl-6-oxo-3-piperidinyl)- (CA INDEX NAME)

IT

06/05/2008

L4 ANSWER 43 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 1973:110501 CAPLUS
OCCUMENT NUMBER: 78:110501
ORIGINAL REFERENCE NO.: 78:17743a,17746a

AUTHOR(S): Nitrile salts. I. Dimerization of nitriles having α-hydrogen in the presence of hydrogen chloride
AUTHOR(S): Kataqiri, Ichiro; Komori, Saburo

CORPORATE SOURCE: Fac. Eng., Osaka Univ., Suita, Japan
Bulletin of the Chemical Society of Japan (1973), 46(1), 292-9

CODEN: BCSJA8; ISSN: 0009-2673

DOCUMENT TYPE: Journal
LANGUAGE: Most were dimers and had the structure investigated. Most were dimers and had the structure
LENH-IC (CHRR2)NRCC1:CRR1.Cl-; hydrolysis gave HM(CONDRR1)2.

IT 40645-76-9P

RI: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
RN 40645-76-9 CAPLUS
CN Benzeneethanimidamide, N-(1-chloro-2-phenyl-1-propen-1-yl)-α-methyl-, hydrochloride (1:1) (CA INDEX NAME)

Ph NH C1 Ph | || | | Me-CH-C-NH-C== C-Me

● HCl

L4 ANSWER 45 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 1973:84378 CAPLUS
COCUMENT NUMBER: 78:84378
CRIGINAL REFERENCE NO: 78:13469a,13472a

Heso, racemic, and optically active forms of 3,6-bis[1-hydroxy-1-(4-methylphenylethyl]-1,2,4,5-tetrazines and related systems along with the corresponding 3,5-disubstituted 1,2,4-triazoles, their

4-amino derivatives, and 2,5-disubstituted 1,3,4-oxadiazoles including their circular dichroism spectra

AUTHOR(S): Neilson, D.G., Mahmood, Safia; Watson, K. M.
Dep. Chem., Univ. Dundee, Dundee, UK
SOURCE: Journal of the Chemical Society, Perkin Transactions 1: Organic and Bio-Organic Chemistry (1972-1999) (1973), (4), 335-9 (COENT. JCPR84, ISSN: 0300-922X Journal
LANGUAGE: English
CTHER SOURCE(S): CASREACT 78:84378
CTHER SOURCE(S): CASREACT 78:84378
CTHER SOURCE(S): Reduction of I gave the corresponding 1,2-dihydrotetrazines
and HZNNH2.H2O. Reduction of I gave the corresponding 1,2-dihydrotetrazines
(II) which rearranged in HCl-MeOH to give 4-amino-1,2,4-triazoles (III). Deamination of III with HNO2 gave 3,5-bis[1-hydroxy-1-(4-methylphenyl)-1-(4)-phenylethyl)-1,2,4-triazoles. A mixture of meso- and (t)-I with MeCO3H gave 2,5-bis[1-hydroxy-1-(4-methylphenyl)-1]-1,3,4-oxadiazole. The 1-hydroxy-1-phenylethyl and 1-hydroxy-1-(4-methylphenyl)-1]-1,3,4-oxadiazole. The 1-hydroxy-1-phenylethyl and 1-hydroxy-1-(4-methylphenyl)-1]-1,3,4-oxadiazole. The 1-hydroxy-1-phenylethyl and 1-hydroxy-1-(4-methylphenyl)-1]-1,3,4-oxadiazole. The 1-hydroxy-1-phenylethyl and 1-hydroxy-1-henylethyl and 1-hydroxy-1-henylethyl and 1-hydroxy-1-(4-methylphenyl)-1]-1,3,4-oxadiazole. The 1-hydroxy-1-phenylethyl and 1-hydroxy-1-hydroxy-1-henylethyl and 1-hydroxy-1-hydroxy-1-henylethyl and 1-hydroxy-1-hydroxy-1-hydroxy-1-henylethyl and 1-hydroxy-1-hydroxy-1-henylethyl and 1-hydroxy-

14 ANSWER 44 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 1973/97594 CAPLUS
DOCUMENT NUMBER: 78:97594
ORIGINAL REFERENCE NO.: 78:15663a,15666a
Nitrile salts. II. Facile one-step synthesis of the pyrimidine nucleus
AUTHOR(S): Yanajda, Shozo, Fujita, Tetsuo; Ohoka, Masataka; Kumaqai, Reiji; Komori, Saburo
CORPORATE SOURCE: Fac. Eng., Osaka Univ., Suita, Japan
SOURCE: Bulletin of the Chemical Society of Japan (1973), 46(1), 299-302
CODEN: BCSJA8; ISSN: 0009-2673
DOCUMENT TYPE: Journal
LANGUAGE: English
GI For diagram(s), see printed CA Issue.
AB The reaction of N-(α-chloroalkenyl)alkylamidine hydrochlorides (I) prepared from nitriles with two α-hydrogens reacted with COC12 at 100-105° to give good yields of 4,6-dichloro-2,5-disubstituted-pyrimidines (II). I, which were obtained from nitriles with only one α-hydrogen, afforded 2-alkylidene-4,6-dichloro-5,5-disubstituted-2,5-dihydropyrimidines (III) in good yields.

IT 40645-76-9
RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, with phosgene, pyrimidines by)
RN 40645-76-9 CAPLUS
CN Benzenethanimidanide, N-(1-chloro-2-phenyl-1-propen-1-yl)-α-methyl-, hydrochloride (1:1) (CA INDEX NAME)

● HCl

L4 ANSWER 45 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) monohydrochloride (9CI) (CA INDEX NAME)

● HCl

RN 941-52-6 CAPLUS
CN Benzeneethanimidamide, α -hydroxy- α , 4-dimethyl-, monohydrochloride, (-)- (9CI) (CA INDEX NAME)

Rotation (-).

● HCl

RN 941-51-5 CAPLUS RN Benzeneethanimidamide, α -hydroxy- α ,4-dimethyl-,

Rotation (+).

● HCl

ANSWER 46 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1971:405728 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 75:5728 75:951a.954a

TITLE:

75:951a,954a

a-Phenyl carboxylic acid compounds
Rossi, Alberto
CIBA Ltd.
Ger. Offen., 98 pp.
CODEN: GWXXBX
Patent PATENT ASSIGNEE(S): DOCUMENT TYPE.

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
DE 2025518	A	19701210	DE 1970-2025518		19700526
CH 559173	A5	19750228	CH 1969-8650		19690605
CH 573909	A5	19760331	CH 1970-6221		19700424
CA 980783	A1	19751230	CA 1970-83721		19700526
US 3801581	A	19740402	US 1970-41107		19700527
ZA 7003642	A	19710127	ZA 1970-3642		19700528
FR 2052932	A1	19710416	FR 1970-20213		19700602
FR 2052932	A5	19710416			
ES 380333	A1	19720916	ES 1970-380333		19700603
BE 751451	A	19701204	BE 1970-751451		19700604
NL 7008158	A	19701208	NL 1970-8158		19700604
GB 1319251	A	19730606	GB 1970-27284		19700605
GB 1319252	A	19730606	GB 1972-55884		19700605
US 3853892	A	19741210	US 1973-338698		19730307
RIORITY APPLN. INFO.:			CH 1969-8650	Α	19690605
			CH 1969-18441	A	19691211
			CH 1970-6221	Α	19700424
			CH 1969-6221	A	19700424

Title compds., useful as anti-inflammatory agents, have the structure ACGHGKNRCX, in which A = azacycloalkyl or -alkenyl, Rl and R2 are H = alkyl, and X = CO2H or a derivative Thus, 4-phenylpjeridine treated that the contract of the cont AB with

C5H5N and AcCl give 1-acetyl-4-phenylpiperidine (I). I, AcCl, and CS2 is treated with AlCl3 to give 1-acetyl-4-(p-acetylphenyl)piperidine (II).

ethanolic

nolic KOH, then HCl gives the HCl salt of $\alpha-\{p-(4-piperidyl)phenyl]propionic acid, which is converted to its Et ester (V), then acetylated to give ethyl <math display="inline">\alpha-[p-(1-acetyl-4-$

US 1970-41107

A2 19700527

L4 ANSMER 47 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1969:42424 CAPLUS

DOCUMENT NUMBER: 70:42424

Optical rotatory dispersion of α-hydroxy
optical rotatory dispersion of α-hydroxy
amidines and their transition metal complexes

Neilson, Douglas G.

CORFORATE SOURCE: Univ. St. Andrews, Dundee, UK
SOURCE: (1967), Meeting Date 1966, 186-91. Editor(s):
Bonnett, R. United Trade Press Ltd.: London, Engl.
CODEN: 20LHA8

CODEN: 20LHA8

O.R.D. of mandelamidinium chlorides (I) and lactamidinium chloride (II)
were measured in MeOH or H2O to obtain their absolute configuration, but

the

were measured in MeOH or H2O to obtain their absolute configuration, but results were rather irregular: no full Cotton effect curves could be measured for (-)-I [R = H, 2-Cl, and 2-Br] and (-)-II, while 2 extrema were observed for (+)-I [2-MeO, 2-EtO, 4-MeO and 4-EtO]. Thus, 0.R.D. of the Cu complexes were measured; all the Cu complexes of α-hydroxyamidines of known D-configuration exhibited a pos. Cotton effect, which permitted the D-configuration to be assigned to I [2-MeO, 2-EtO, 2-Cl, 2-Br, 4-MeO, 4-EtO, 3-EtO and 2,4-di-Cl] for which chemical methods cannot be applied owing to the facile racemization. The Cu complex of D-(+)-II gave a pos. 0.R.D. curve, establishing the greater value of 0.R.D. curves of Cu complexes over that of the parent amidines for the correlation of configuration. The Ni complex is also effective but proved difficult to synthesize. 0.R.D. curves of some of the Cu complexes of I [2-EtO, 3-EtO and 4-MeO] have an addnl. extrema near 270 mμ. The Cotton effect owing to the ligand is counterbalanced by an effect of opposite sign but approx. equal intensity owing to the complex as a whole. Support for this argument was given by comparing the ular circular

dichroism curves of I [2-C1] and I [2-Et0] and their Cu complexes. O. R. D.

D.

of compds. containing the amidine group in a heterocyclic ring (e.g., imidazoline) are also discussed.

22210-97-5
RL: FRCC (Process)
 (optical rotary dispersion of)
22210-97-5 CAPLUS

Mandelamidine, α-methyl-, monohydrochloride, (+)- (8CI) (CA INDEX NAME)

Absolute stereochemistry.

HCl

ANSWER 46 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) piperidyl)phenyl]propionate, hydrolysis of which gives α-[p-(1-acety1-4-piperidyl)phenyl]propionic acid. IV was similarly prepd. from 2-(p-bromophenyl)-2-methyl-1, 3-dioxolane Grignard reagent and 1-benzy1-4-piperidone via 2-[p-(1-benzy1-4-hydroxy-4-piperidyl)phenyl]-2-methyl-1, 3-dioxolane, p-(1-benzy1-1, 2,5,6-tetrahydro-4-pyridyl)acetophenone, and 1-hydroxy-1-[p-(4-piperidyl)phenyl]ethane. 1-Methyl-2-coxo-5-[p-(1-cyanoethyl)phenyl]piperidine is treated with NH2OH.HCl to give α-[p-(1-methyl-2-oxo-5-piperidyl)phenyl]phenyl]phenyl]phenylphen acid morpholide, which is hydrolyzed to p-(4-piperidyl)phenylacetic acid-HCl.
32262-02-5P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
32262-02-5 CAPLUS
Hydratropamidine, N-hydroxy-p-(1-methyl-6-oxo-3-piperidyl)- (8CI) (CA
INDEX NAME)

L4 ANSWER 47 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

ANSWER 48 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1965:454040 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 63:54040 63:9784a-c

63:9784a-c
Optical rotatory dispersion. XIX. A series of acids,
imidazolines, amidinium chlorides, and their copper
complexes, related to mandelic acid
Emerson, T. R.; Ewing, D. F.; Klyne, W.; Neilson, D.
G.; Peters, D. A. V.; Roach, L. H.; Swan, R. J.
Univ. London, Swed.
Journal of the Chemical Society (1965), (July),
4007-14
COMPN. ICCOMP. ICCOMP. 155N. 0368-1568 TITLE:

AUTHOR(S):

CORPORATE SOURCE:

SOURCE:

Journal of the Chemical Society (1965), (July), 4007-14

CODEN: JCSOA9; ISSN: 0368-1769

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB The optical rotatory dispersion (o.r.d.) curves of series of α-hydroxy acids related to mandelic acid show that the Cotton-effect curves observed are generally due to the n + π* transition of the carboxyl group and not to the phenyl absorption band (260-280 mμ). The o.r.d, curves for the related amidinium chlorides show distinct extrema in the 250-280 mμ region when the phenyl group carries an alkoxy-substituent. The o.r.d. curves of the amidinium chlorides, however, are more complex than those of their parent acids and not so useful for configurational assignments. Cu complexes derived from these α-hydroxyamidnium chlorides show a Cotton effect at .appxx.590 mμ. Compds of D-configuration have a positive Cotton effect in this region. This rule has permitted the assignment of configuration to some 10 amidines, not previously correlated by chemical means.

IT 941-52-6 92442-87-0

(Derived from data in the 7th Collective Formula Index (1962-1966))

RN 941-52-6 CAPLUS

Benzeneethanimidamide, α-hydroxy-α,4-dimethyl-, monohydrochloride, (-)- (9CI) (CA INDEX NAME)

Rotation (-).

• HCl

92442-87-0 CAPLUS Benzeneethanimidamide, α -hydroxy- α -methyl-, monohydrochloride (9C1) (CA INDEX NAME)

ANSWER 48 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) Mandelamidine, m, α -dimethyl-, hydrochloride (7CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{Me} & \text{NH} \\ & \parallel & \parallel \\ \text{C-C-NH}_2 \\ & \text{OH} \end{array}$$

L4 ANSWER 48 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

● HCl

941-51-5, Mandelamidine, p, α -dimethyl-, hydrochloride, D-(-)- 4023-95-4, Mandelamidine, α -methyl-, hydrochloride, D-(-)- 94281-37-5, Mandelamidine, m, α -dimethyl-, hydrochloride, D-(-)- (optical rotatory dispersion and spectrum of) 941-51-5 CAPLUS Benzeneethanimidamide, α -hydroxy- α , 4-dimethyl-, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

4023-95-4 CAPLUS Mandelamidine, $\alpha\text{-methyl-, hydrochloride, D-(-)- (8CI)}$ (CA INDEX NAME)

Absolute stereochemistry.

● HCl

RN 94281-37-5 CAPLUS

L4 ANSWER 49 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 1965:454039 CAPLUS
OCCUMENT NUMBER: 63:54039
ORIGINAL REFERENCE NO.: 63:9783h, 9784a
TITLE: Optical rotatory dispersion. XV. Monosubstituted succinic acids
AUTHOR(S): Fredga, A.; Jennings, J. P.; Klyne, W.; Scopes, Patricia M.; Sjoberg, B.; Sjoberg, S.
Univ. Uppsala, Swed.
SOURCE: Journal of the Chemical Society (1965), (July), 3928-33
CODEN: JCSOA9; ISSN: 0368-1769
DOCUMENT TYPE: Journal
LANSUAGE: English
AB of. CA 62, 13191b; 63, 7065g. The ORD curves of many α-substituted succinic acids are measured. All these compds. show Cotton effects associated with the carboxyl absorption band at about 225 mμ.
α-Alkyl-, α-aryl-, and α-halosuccinic acids of the D-configuration all give pos. Cotton effects in water and in MeOH; D-α-Alkylthiosuccinic acids give somewhat more complex pos. curves. D-α-Hydroxysuccinic acid (D-malic acid) and its O-alkyl ethers give neg. Cotton effects in water and in MeOH. The signs of the dispersion curves of most of these acids are reversed on the addition of alkali.
IT 941-51-5 941-52-6 92442-87-0
(Derived from data in the 7th Collective Formula Index (1962-1966))
RN 941-51-5 CAPLUS
CN Benzeneethanimidamide, α-hydroxy-α, 4-dimethyl-, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

941-52-6 CAPLUS Benzeneethanimidamide, α -hydroxy- α ,4-dimethyl-, monohydrochloride, (-)- (9CI) (CA INDEX NAME)

Rotation (-).

L4 ANSWER 49 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

● HCl

92442-87-0 CAPLUS Benzeneethanimidamide, α -hydroxy- α -methyl-, monohydrochloride (9CI) (CA INDEX NAME)

ANSWER 50 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) 3-(p-chlorobenzyl)-5-(β -morpholinoethyl)-1,2,4-oxadiazole-HCl, m. 179-80°, 3-(p-chlorobenzyl)-5-piperidinomethyl)-1,2,4-oxadiazole-HCl, m. 157°, 3-(p-chlorobenzyl)-5-piperidinomethyl)-1,2,4-oxadiazole-HCl, m. 155°, 3-[β -phis(4-chlorophenyl)ethyl]-5-piperidinomethyl)-1,2,4-oxadiazole-HCl, m. 155°, 3-[β -phis(4-chlorophenyl)ethyl]-5-piperidinomethyl)-1,2,4-oxadiazole H maleate, m. 117°, 3-(β -pibis(4-chlorophenyl)hyl)-5-[β -piperidinomethyl)-1,2,4-oxadiazole H maleate, m. 126°, 3-[bis(3,4-dimethoxyphenyl)methyl)-5-[β -morpholinomethyl)-1,2,4-oxadiazole, m. 112-13° (abs. EtOH), 3-(α , β -diphenylethyl)-5-[β -[β -piperidinomethyl)-1,2,4-oxadiazole-BCl, m. 186-7° (abs. EtOH), 3-(α , β -diphenylethyl)-5-[β -(N-piperidinomethyl)-1,2,4-oxadiazole-BCl, m. 191° (96% EtOH), 3-(β - β -diphenylethyl)-5-[β -(N-methylpiperazino)ethyl]-1,2,4-oxadiazole, m. 83° [dihydrochloride m. 205-7° (96% EtOH)], 3-diphenylmethyl-5-[β -(N-methylpiperazino)ethyl]-1,2,4-oxadiazole, m. 37° [dihydrochloride m. 196° (96% EtOH)], 3-(β -chlorobenzyl)-5-[β -(N-methylpiperazino)ethyl]-1,2,4-oxadiazole-2BCl, m. 186.5-8.5° (96% EtOH), 3-(β - β -(β -N-methylpiperazino)ethyl]-1,2,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β -diphenylmethyl)-1,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β -diphenylmethyl)-1,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β -diphenylmethyl)-1,2,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β -diphenylmethyl)-1,2,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β -diphenylmethyl)-1,2,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β -diphenylmethyl)-1,2,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β -diphenylmethyl)-1,2,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β -diphenylmethyl)-1,2,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β - β -diphenylmethyl)-1,2,4-oxadiazole-2BCl, m. 191-3° (96% EtOH), and 3-(β - β - β -diphenylmethylmethyl)-1,2,4-oxadiazole-2BCl, m. 191-3° (96% Et

24 hrs. and filtered to give 44.59 g. O-y-chlorobutyryl- β,β -diphenylpropionamidoxime (II), m. 145°. II (10.35 g.) and Ac20 (6 ml.) heated on a water-bath, treated with H2O and C6H6, and the org. phase washed with Na2CO3 soln. and concd. gives 9.90 g. $3-(\beta,\beta$ -diphenylethyl)-5-(y-chloropropyl)-1,2,4-oxadiazole (III) (m.p. not given). A mixt. of 9.90 g. III, 45 ml. PhMe, and 8.90

[III] (m.p. not given). A mixt. of 9.90 g. III, 45 ml. PhMe, and 8.90 piperidine is refluxed 11 hrs. and filtered and the filtrate washed with H2O and evapd. to yield 11.20 g. 3-(β,β-diphenylethyl)-5(γ-piperidinopropyl)-1,2,4-oxadiazole. H maleate m. 146-79
(Me2CO). The γ-morpholinopropyl deriv. is prepd. similarly, H maleate m. 133°. A solm. of 14.53 g. β-chloropropionyl chloride in Me2CO is added dropwise with stirring at 0-5° to a suspension of 27.6 g. 1 and 9.86 g. NaHCO3 in 140 ml. abs. Me2CO and the whole stirred 7 hrs. and added to 1100 ml. H2O to yield 27.8 g. O-(β-chloropropionyl)-β,β-diphenylpropionamidoxime (IV), m. 116-17° (abs. EtOH or C6H6). A solm. of 4.55 g. IV in 25 ml. abs. PhMe and 3 ml. piperidine is refluxed 7 hrs., 20 ml. H2O added, the mixt. evapd. to dryness in vacuo, and the residue treated with HCl in EtOH to give 2.15 g. 3-(β,β-diphenylethyl)-5-(β-piperidinoethyl)-1,2,4-oxadiazole-HCl, m. 192° (abs. EtOH). O-Chloroacetyl-β,β-diphenylethyl)-5-chloromethyl-1,2,4-oxadiazole, m. 73-4° (McOH). Heating it with piperidine in PhMe and treating the product with HCl in EtOH to ppt. 3-(β,β-diphenylethyl)-5-(β-piperidinoethyl)-1,2,4-oxadiacole, hCl, and labs. C5HSN at 20°, the mixt. refluxed 2 hrs. and evapd. to dryness in v

ANSWER 50 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1965:66553 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 62:66553 62:11821e-h.11822a-e 62:11821e-h,1822a-e
1,2,4-Oxadiazoles with pharmaceutical effect
Harsanyi, Kalman; Kiss, Pal; Korbonits, Dezso;
Malyata, Ilona; Erdelyi, Ilona; Tardos, Laszlo;
Leszkowszky, Gyorgy
Chinoin Gyogyszer es Vegyeszeti Termekek Gyara Rt.
23 pp.
Patent INVENTOR(S): PATENT ASSIGNEE(S): DOCUMENT TYPE: LANGHAGE . Unavailable FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND APPLICATION NO. DATE DATE HU 151748 BE 645822 NL 302339 US 3280122 PRIORITY APPLN. INFO.: 19641223 19630329 NL US 1964-354465 HU 19661018 19640324 19630329

OTHER SOURCE(S): MARPAT 62:66553
AB A mixture of 24 g. β,β-diphenylpropionamidoxime(I), 37 g. Et
β-piperidinopropionate, 200 ml. absolute EtOH, and 2.3 g. Na is refluxed
8 hrs., concentrated in vacuo, 200 ml. H2O and 4.0 g. NaOH are added,

mixture is extracted with C6H6. The organic phase is concentrated in

L4 ANSWER 50 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) oxadiazole-2HCl, m. 192-3°. These products showed spasmolytic, local anesthetic, cough-reliever, analgesic, anti-inflammatory, antipyretic, and circulation influencing effects.

IT 988-45-6P, Propionamidoxime, 2,2-diphenyl-, O-chloroacetate 971-95-9P, Propionamidoxime, 2,2-diphenyl-, O-3-chloropropionate 974-34-SP, Propionamidoxime, 2,2-diphenyl-, O-4-chlorobutyrate R1: PREP (Preparation) (preparation of)
RN 988-45-6 CAPLUS
ON Propionamidoxime, O-(2-chloroacetyl)-2,2-diphenyl-, O-chloroacetate (8CI)

(CA INDEX NAME)

971-95-9 Propionamidoxime, 2,2-diphenyl-, O-(3-chloropropionyl)- (8CI) (CA INDEX NAME) CAPLUS

$$\begin{array}{c|c} & & \text{HN Ph} \\ \parallel & \parallel & \parallel \\ \text{ClCH}_2-\text{CH}_2-\text{C-O-NH-C-C-Me} \\ & \parallel & \parallel \\ & \text{Ph} \end{array}$$

974-34-5 CAPLUS Propionamidoxime, O-(4-chlorobutyry1)-2,2-dipheny1- (8CI) (CA INDEX

10/568,760

Page 33

L4 ANSWER 51 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1965:36364 CAPLUS

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.:

TITLE:

AUTHOR(S): CORPORATE SOURCE:

1965:36364 CAPLUS
62:36364
62:6374f
The resolution of some substituted lactamidines and atrolactamidines by means of the mandelic acids Ewing, D. F.; Neilson, D. G. Univ. St. Andrews, Dundee, UK
Journal of the Chemical Society (1965), (Jan.), 770-4
CODEN: JCSOA9; ISSN: 0368-1769
Journal SOURCE:

DOCUMENT TYPE:

DOCUMENT TYPE: Journal
LANGUAGE: English

AB m- and p-Methylatrolactamidines were prepared from the corresponding methylacetophenones and were resolved by means of the mandelic acids. o-Methylacetophenone failed to give an amidine.

α-Benzyl-lactamidine was also resolved by means of these acids but Pphenyllactamidine showed no separation of the diastereoisomers.

IT 941-50-4 941-52-6 943-23-7 971-52-8 971-53-9 (Derived from data in the 7th Collective Formula Index (1962-1966))

RN 941-50-4 CAPLUS

CN Benzeneethanimidamide, α-hydroxy-α,4-dimethyl-, monohydrochloride, (+)- (9CI) (CA INDEX NAME)

941-52-6 CAPLUS Benzeneethanimidamide, α -hydroxy- α , 4-dimethyl-, monohydrochloride, (-)- (9CI) (CA INDEX NAME)

Rotation (-).

L4 ANSWER 51 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

HC1

943-23-7 CAPLUS Mandelamidine, m, α -dimethyl-, monohydrochloride, (-)- (8CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

971-52-8 CAPLUS Mandelic acid, (S)-, compd. with (-)- α -hydroxy-m-methylhydratropamidine (1:1) (8CI) (CA INDEX NAME)

Rotation (-).

CRN 53623-24-8 CMF C10 H14 N2 O

L4 ANSWER 51 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

Absolute stereochemistry. Rotation (+).

971-53-9 CAPLUS Mandelic acid, (R)-, compd. with (+)-p, α -dimethylmandelamidine (1:1) (8c1) (CA INDEX NAME)

CM 1

CRN 46147-67-5 CMF C10 H14 N2 O

Rotation (+).

CM 2

Habte

Absolute stereochemistry. Rotation (-).

L4 ANSWER 51 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

109595-37-1, Mandelamidine, α -methyl- (derivs., resolution by mandelic acids) 109595-37-1 CAPLUS Benzeneethanimidamide, α -hydroxy- α -methyl- (CA INDEX NAME)

941-51-5P, Mandelamidine, p,α-dimethyl-, hydrochloride, isomers 94281-37-5P, Mandelamidine, m,α-dimethyl-, hydrochloride, isomers 95157-76-9P, Mandelic acid, compound with m,α-dimethylmandelamidine (1:1), (+) = 95157-78-1P, Mandelic acid, compound with p,α-dimethylmandelamidine (1:1), isomers RL: PREP (Preparation) (preparation of) 941-51-5 CAPLUS Benzeneethanimidamide, α-hydroxy-α,4-dimethyl-, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

94281-37-5 CAPLUS Mandelamidine, m, \alpha-dimethyl-, hydrochloride (7CI) (CA INDEX NAME)

L4 ANSWER 51 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

HCl

95157-76-9 CAPLUS Mandelic acid, compd. with $m,\alpha\text{-dimethylmandelamidine}$ (7CI) (CA INDEX NAME)

CM

CMF C8 H8 O3

Ph но— сн— со₂н

95157-78-1 CAPLUS Mandelic acid, compd. with p, α -dimethylmandelamidine (7CI) (CA INDEX NAME)

CRN 95157-77-0 CMF C10 H14 N2 O

L4 ANSWER 51 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

CM 2

Рh | но— сн— со₂н

L4 ANSWER 52 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 1965:36363 CAPLUS
COUMENT NUMBER: 62:36363
CRIGINAL REFERENCE NO.: 62:6374c-f
TITLE: Electrophilic substitution at saturated carbon. XXIV.
Trifluoromethyl as a carbanion-stabilizing group
Cram, Donald J.; Wingrove, Alan S.
SOURCE: Journal of the American Chemical Society (1964),
86(24), 5490-6
CODEN: JACSSAT; ISSN: 0002-7863
DOCUMENT TYPE: Journal
LANCUAGE: English
AB Two systems have been prepared for study of the stereochem. course of the base-catalyzed H-D exchange at C attached to a trifluoromethyl group.
Optically active 2-methyl-3-phenyl-1,1,1-trifluoropropane (I) and the

compound deuterated in the 2-position, and optically active 2-phenyl-1,1,1-trifluorobutane (II) and its deuterated counterpart (2-position) were examined In tert-BuOD at 124°, (+)-I was found to undergo elimination reaction to the exclusion of isotopic exchange. The initially formed 1,1-difluoro-2-methyl-3-phenyl-1-propene underwent a base-catalyzed allylic rearrangement to give a 6.5:1 mixture of trans-to cis-3,3-difluoro-2-methyl-1-phenyl-1-propene (trans-to cis-III), which were identified by their spectral properties. The base-catalyzed elimination reaction exhibited a kinetic isotope effect of 1.2, a fact which suggests a carbanion intermediate for the reaction. II also underwent elimination to give 1,1-difluoro-2-phenyl-1-butene and its polymers. However, H-D exchange also occurred, but at a much slower

In tert-BuOH-tert-BuOK, and in EtOH-KOEt, isotopic exchange went with total racemization ($ke/k\alpha$, the ratio of the rate constant for exchange to the rate constant for racemization, was equal to unity). In

to the rate constant for racemization, was equal to unity). In MeOH-KCMP, when the control of th

9915/-/8-1 (Derived from data in the 7th Collective Formula Index (1962-1966)) 941-50-4 CAPLUS

Senzeneethanimidamide, α-hydroxy-α,4-dimethyl-, monohydrochloride, (+)- (9CI) (CA INDEX NAME)

Rotation (+).

■ HC1

(Continued) L4 ANSWER 52 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN

941-51-5 CAPLUS Benzeneethanimidamide, $\alpha\text{-hydroxy-}\alpha,4\text{-dimethyl-,}$ monohydrochloride (9CI) (CA INDEX NAME)

941-52-6 CAPLUS Benzeneethanimidamide, α -hydroxy- α , 4-dimethyl-, monohydrochloride, (-)- (9CI) (CA INDEX NAME)

• HCl

943-23-7 CAPLUS

Mandelamidne, m, α -dimethyl-, monohydrochloride, (-)- (8CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

L4 ANSWER 52 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) 971-52-8 CAPLUS Mandelic acid, (S)-, compd. with (-)- α -hydroxy-m-methylhydratropamidine (1:1) (8CI) (CA INDEX NAME) CRN 53623-24-8 CMF C10 H14 N2 O Rotation (-).

Absolute stereochemistry. Rotation (+).

971-53-9 CAPLUS Mandelle acid, (R)-, compd. with (+)-p, α -dimethylmandelamidine (1:1) (8C1) (CA INDEX NAME)

CM 1 CRN 46147-67-5 CMF C10 H14 N2 O

Rotation (+).

ANSWER 52 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

CM 2

но— сн— со2 н

L4 ANSWER 52 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

CM 2

Absolute stereochemistry. Rotation (-).

94281-37-5 CAPLUS Mandelamidine, m, α -dimethyl-, hydrochloride (7CI) (CA INDEX NAME)

● HCl

95157-78-1 CAPLUS Mandelic acid, compd. with p, α -dimethylmandelamidine (7CI) (CA INDEX NAME)

CM 1

CRN 95157-77-0 CMF C10 H14 N2 O

L4 ANSWER 53 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 1962:435850 CAPLUS
DOCUMENT NUMBER: 57:35850
ORIGINAL REFERENCE NO.: 57:7081i, 7082a-d
TITLE: The structure of N-mono- and N,N'-disubstituted anidines
AUTHOR(S): Prevorsek, Dusan C. Textile Res. Inst., Princeton, NJ
SOURCE: Journal of Physical Chemistry (1962), 66, 769-78
CODENT TYPE: Journal OF Physical Chemistry (1962), 66, 769-78
CODENT TYPE: Journal Unavailable
AB Factors influencing the position of tautomeric equilibrium of a number of N-mono- and N,N'-disubstituted amidines were studied by infrared spectral analyses. In solution the unsubstituted amidines, RC(:NR:NH2.dblarw.
RC(:NN)MHR' (I) (R' = H), existed as a mixture of approx. equal amts. of each tautomer. The equilibrium of I was found to be displaced in proportion to
the electroneg. of the substituents R'. Thus, when R' was phenyl or hydroxyl (amidoximes), the equilibrium was shifted to a the left, whereas an anyment of the R.

whereas

ethyl group shifted the equilibrium to the right. The nature of the R group

apparently was without effect. Characteristic frequency assignments in the 2-7° region for eight N-monosubstituted amidines and seven amidoximes were given where R varied from 2-thienyl, 2-, 3-, or 4-piperidyl, benzyl, α-phenethyl, α-phenylpropyl, and αphenylbutyl groups, R' = H, and R' = hydroxyl, phenyl, methyl, or ethyl. The spectra of N,N'-disubstituted amidines (II) in dilute tion

solution ion showed two bands in the 3 μ region, b suggesting the presence of either two forms of a monomer or a single form giving rise to both bands. Geometric isomerism with respect to the C:N bond was felt unlikely

of the steric effects offered by the R' and R'' groups (substituted phenvl

or naphthyl groups). The possibility that one band was an overtone of

fundamental C:N stretching vibration in the 6 μ region was also deemed improbable. Simple tautomerism could not explain the two bands, since identical configurations would result when R' = R''. It was concluded, however, that N,N'-disubstituted amidines very probably exhibited in solution

tautomerism leading to a rotational isomerism with respect to both single and double CN bonds. This would explain the appearance of two N-N and

and double CN bonds. This would explain the appearance of two N-N and bands for derivs. with identical substituents. Characteristic frequency assignments in the 2-7 μ region for ten N,N'disubstituted amidines were given where R = methyl, α -phenethyl, and α -phenylpropyl, R' and (or) R'' = ethyl, phenyl, substituted phenyl, or β -naphthyl. The infrared spectra of these N-mono- and N,N'-disubstituted amidines d indicated an electronic configuration similar to that of amides. All the amidines studied were prepared according to known procedures. 91429-53-7P, Hydratropamidine, N-ethyl- 92579-12-9P, Hydratropamidine, N-phenyl-RL: PREP (Preparation) (preparation of) 91429-53-7 CAPLUS Hydratropamidine, N-ethyl- (6CI, 7CI) (CA INDEX NAME)

L4 ANSWER 53 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

Ph NH | || Me-CH-C-NHEt

92579-12-9 CAPLUS Hydratropamidine, N-phenyl- (6CI, 7CI) (CA INDEX NAME)

HN Ph || | | PhNH-C-CH-Me

ANSWER 54 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) converted by SOC12 into 96% PhMeCHCOCI (VII), b13 100-1°. The following PhCHRCOCH and chlorides were prepd. (R, m.p., b.p./mm., and % yield of acid, and b.p./mm. and % yield of chloride given): H, 78°, 144°/12. 92, 95°/14, 88; Me, about 16°, 145°/13, 89, 110°/13, 96; Et., 42°, 158°/15, 85, 112°/16, 97, Pr. 52% 167°/15, 90, 118°/13, 96; n-C8H17, -, 170-3°/0.3, 93, 138°/0.3, 88. The various amides were prepd. from the COC1 derivs; e.g., 8.5 g. VII and 5 g. EtNH2 were each dissolved in 75 cc. C6H6, stirred together, and refluxed 0.5

were each dissolved in 75 cc. C6H6, stirred together, and refluxed 0.5 (1 hr. for the higher homologs). After cooling, EtNB2.HCl was dissolved in H2O, the soln. extd. with E2O and the ext. added to the C6H5 layer, dried over Na2SO4, and the solvents removed in vacuo. The residue of PhMcCHCONHET crystd. on cooling; yield after 2 crystns. from C6H6-petr. ether (6:92) 83%, m. 65.5-6*. PhMcCHCONH2 was prepd. similarly, using excess NH3 and not allowing the temp. to exceed 65°, yield 83% after crystn. from C6H6. m. 92.5°. The following PhcHRCONN'R'' were prepd. (R, R', R' '157°, H, H, Et, 96, -, 69, 5°; H, Et, Et, 98, 113°/0.3, -; H, H, Ph, 95, -, 113°; H, H, PhCH2, 93, -, 119°; Me, H, H, 89, -, 92.5°, Me, H, Et, 95, -, 66°; Me, Et, Et, 94, 105°/0.15, about 16°; Me, H, Ph, 94, -, 134°; Me, H, FhCH2, 96, -, 75°; Et, H, H, 97, -, 84°, Et, H, Et, 93, -, 66°; Et, Et, 96, 115°/0.2, 22°, 5; Et, H, Ph, 96, -, 97°; Et, H, PhCH2, 93, -, 82°; n-C8H17, H, H, 99, -, 86°; n-C8H17, H, Et, 95, -, 53°; n-C8H17, H, PhCH2, 99, -, 52°. R662-24-9P, Hydratropamidine, N-phenyl-

N-phenyl-RL: PREP (Preparation)

(preparation of) 78622-24-9 CAPLUS

Benzeneethanimidamide, α-methyl-, monohydrochloride (9CI) (CA INDEX

91429-53-7 CAPLUS Hydratropamidine, N-ethyl- (6CI, 7CI) (CA INDEX NAME)

Ph NH | | | | Me-CH-C-NHEt

92579-12-9 CAPLUS Hydratropamidine, N-phenyl- (6CI, 7CI) (CA INDEX NAME)

ANSWER 54 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN SSION NUMBER: 1962:410679 CAPLUS MENT NUMBER: 57:10679
INAL REFERENCE NO.: 57:2141d-i,2142a-b ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.:

HINAL REFERENCE NO.: 57:2141d-i,2142a-b
E: Amidines and other derivs. of phenylalkylacetic acids
OR(S): Delaby, Raymond; Reynaud, Pierre; Lilly, Franck
CE: Bulletin de la Societe Chimique de France (1961) 2067
CODEN: BSCFAS; ISSN: 0037-8968
MENT TYPE: Journal
UNAGE: Unavailable
R SOURCE(S): CASEBACT 57:10679
Starting with phenylalkylacetonitriles, the title compds. were prepared TITLE: AUTHOR(S): SOURCE: DOCUMENT TYPE:

OTHER SOURCE(S):

AB Starting with phenylalkylacetonitriles, the title compds. were prepared in order to ascertain their possible hypocholesterolemic action (Redel and Cottet, CA 48, 13061; C., et al., 13975c). The first step was the preparation of the HCl salt of the imino ester; e.g., PhMeCHC(:NH)ORT.HCl (I) was prepared by passing a current of HCl into a mixture of 20 g. PhMeCHCN (II), 40 cc. Et2O, and 40 cc. absolute EtOH at 0-5° 2.5 hrs. and keeping 2 days at 0°. After removing the solvent in vacuo the residual oil was crystallized from Et2O to yield 92% I, m. 106°; free imino ester (III) bi3 116°, n23D 1.5064. PhMeCHC(:NH)NH2.HCl (IV) resulted when 10 g. I in 40 cc. EtOH was treated with NH3 0.5 hr., then refluxed 0.5 hr., the solvent distilled in vacuo, and the residue crystallized from hot H2O, yield, 8.5 g. IV, m. 235°. The following PhCHRC(:NH)OET.HCl were obtained (R, % yield, m.p., b.p., b.p. pressure (mm.) given): Me, 92,106°,--,: Et, 87, 98°, -, -; Pr, 83, 82°, -, The data for PhCHRC(:NH)DCT were Me, 82, -, 116°, 13; Et, 84, -, 121°, 15; Pr, 81, -, 103°, 1. The data for PhCHRC(:NH)DCT were Me, 98, -, 116°, 13; Et, 84, -, -, -, Pr, 99, 238°, -, -. Mono- and dialkylamidines were obtained by the action of Alcl3 (V) on a mixture of nitrile and amine. The following PhCHRC(:NH)NE'R'' HCl were prepared (R, R', R'', % yield, b.p./mm., and

following PhCHRC(:NH)NR'R''.HCl were prepared (R, R', R'', % yield, b.p./mm., and

given): H, H, Et, 85, 110°/0.1, 61°; H, Et, Et, 68, 131°/2.5, -; H, H, Ph, 95, -, 138°; Me, H, Et, 75, 109°/0.2, -; Me, Et, Et, 62, 111°/0.1, -; Me, H, Ph, 84, -, 89°; Et, H, Et, 99, 115°/0.15, -, Et, Et, 50, -, 45°; Et, H, Ph, 91, -, 86°; Pr, H, Et, 67, 152°/3, -; Pr, Et, Et, 54, 102°/0.01, -; Pr, H, Ph, 96, -, 110.5°; n-08H17, H, Et, 75, 143°/0.1, -; n-08H17, Et, Et, 53, 102° n-08H7, H, Et, 75, 143°/0.1, -; n-08H17, Et, Et, 53, 100°/0.3, -; n-08H7, H, Ph, 62, -, 52°. The acids were prepared by saponification of the nitriles. E.g., a solution of 86 g. and 50.3 g.

II in 400 cc. EtOH was refluxed 16 hrs. (NH3 evolution was virtually complete in 12 hrs.), the EtOH distilled in vacuo, the residue dissolved KOH and

H2O and extracted with Et2O to remove neutral compds. (less than 0.2

H2O layer acidified, and the free acid extracted from it with Et2O; yield, 89% PhMe-CHCO2H (VI), b13 145°, m. about 16%, n24D 1.5210. VI was

L4 ANSWER 54 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

HN Ph || | PhNH-C-CH-Me

L4 ANSWER 57 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1959:82995 CAPLUS DOCUMENT NUMBER: 53:82995 CAPLUS SIGNAL REFERENCE NO.: 53:149291,14930a-e Stereochemical structure. XII. Resolution of (1)-atrolactamidinium chloride (ROPER CORPORATE SOURCE: Stereochemical structure. XII. Resolution of (2)-atrolactamidinium chloride (ROPER COPER CORPORATE SOURCE: Queen's Coll., Dundee, UK GURCE: CORPORATE SOURCE: Queen's Coll., Dundee, UK GURCE: CORPORATE SOURCE: Gueen's Coll., Dundee, UK GURCE: GUEEN'S JOURNAL OF THE COLLING (1)-Atrolactamidinium chloride (I) was prepared from PhAc cyanohydrin (II) via Et (1)-atrolactimidate-HCl (III). (1)-Atrolactamidine (IV) was resolved by separation of the diastereoisomeric salts with optically active mandelic acid (V). (-)-Atrolactic acid (VI), isolated from (-)-atrolactamidinium chloride (VII), was of at least 90% optical purity. PhAc (120 g.) in 90 ml. Bt20 and 123 g. NaCN in 150 ml. H20 treated at 5° during 2 hrs. with 210 ml. concentrated HCl, the Ft20 layer separated and the aqueous layer again extracted with Et20, and the ethereal exts. distilled gave 48 g. II, bl8 147-9°, yellow oil. II (48 g.) and 16 g. anhydrous alc. treated 48 hrs. at 0° with 15.2 g. dry HCl and Et20 gave 60 g. III, m. 101-2° (decomposition). III (5 g.) treated with 12 ml. 4N NaOH gave 2 g. Et (:)-atrolactimidate, m. 56-7° (ligroine). An anhydrous solution of 8.5 g. NH3 in 100 ml. alc. shaken 174-5° (ddilute HCl). I (6 g.) shaken at 0° with 15 ml. 10N NaOH and H20 added gave 3.7 g. IV, m. 77-8° (decomposition); picrate m. 188-9°. I (2.5 g.) heated with 2.2 g. Na salt V in H20 to a clear solution gave 1 g. (:)-atrolactamidine (:)-mandelate (IVII), m. 115-9° (H20). I (6.7 g.) and 5.8 g. Na (+)-mandelate heated in 37 ml. H20 gave 2 g. (-)-atrolactamidine (+)-mandelate (HIX), m. 165° (decomposition), [a]165461 12.1° (c 0.91, McOH). Ethereal (+)-mandelate (X) was prepared as in the above method but with (-)-mandelate (X) was prepared as in the above method but with (-)-mandelate (X) was prepared

DOCUMENT NUMBER: 55:47579
CORTIGINAL REFERENCE NO.: 55:91401,9141a
TITLE: Complexes formed by α-hydroxy amidines with transition metal ions
AUTHOR(S): Gould, R. O.; Jameson, R. F.; Neilson, D. G.
COMPORATE SOURCE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT TYPE: Journal
LANGUAGE: Proc. Chem. Soc. (1960) 314-15
DOCUMENT

L4 ANSWER 56 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1961:47579 CAPLUS

ANSWER 57 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

RN 92442-87-0 CAPLUS
CN Benzeneethanimidamide, α-hydroxy-α-methyl-, monohydrochloride

(9CI) (CA INDEX NAME)

Ph NH

Me C C NH2
OH

OH

CN 109595-38-2 CAPLUS
CN Mandelic acid, salt with atrolactamidine (6CI) (CA INDEX NAME)

CM 1

CRN 109595-37-1

CMF C9 H12 N2 O

Ph NH

Me C C C NH2
OH

CM 2

CRN 90-64-2

CMF C8 H8 O3

Ph

HC CH CCH CO2H

IT 109595-37-1, Atrolactamidine, (-)
(and derivs.)
RN 109595-37-1 CAPLUS
CN Benzeneethanimidamide, α-hydroxy-α-methyl- (CA INDEX NAME)

L4 ANSWER 57 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

L4 ANSWER 58 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued)

Рh | но— сн— со₂ н ACCESSION NUMBER: 1958:113428 CAPLUS
DOCUMENT NUMBER: 52:113428
DOCUMENT NUMBER: 52:113428
DOCUMENT NUMBER: 52:113428
DOCUMENT NUMBER: 52:113428

ORIGINAL REFERENCE NO.: 52:2024g-i,20025a

TITLE: from substituted phenylacetic acids
AUTHOR(S): Delaby, Raymond; Reynaud, Pierre; Lilly, Frank
SOURCE: Compt. rend. (1958), 246, 2905-6

DOCUMENT TYPE: Journal
LANSUAGE: Unavailable

AB PHCH2CN with Et2CO3 in the presence of EtCNa gives Et αcyanophenylacetate, converted by treatment with alkyl halides (RX) and
saponification with NaCH to PhCHECN, e.g. 62% PhCHMcCN, b15 108°, 69%
PHCHECN, b15 115° 72% PhCHPTN, b14 130°, and 63%
PHCH(CSH17-n)CN, b0.1 133°. On passing dry HCl into solns. of the
nitriles in EtCH, the iminoesters are formed, and addition of amines in

the

PRESENCE ALCLA GIVEN N-substituted amidines. Thus, PhCHMCCN with HCl
and EtCH gives PhCHMcC(:NH)CEt.HCl, m. 103.5°, and then
PhCHMcH(:NH)NH2.HCl, m. 235°, is ethylated to PhCHMcC(:NH)NHE2.HCl,
b0.2 109°, and PhCHMcC(:NH)CEt.HCl, m. 98° then PhCHEC(:NH)NH2.HCl,
phCHEC(:NH)CHE.HCl, m. 98° then PhCHEC(:NH)NH2.HCl, 322°,
phCHEC(:NH)CET.HCl, m. 88° then PhCHEC(:NH)NH2.HCl, m.
236°, PhCHEC(:NH)NHEP, m. 86°. Alao, PhCHENTOR gives
PhCHETC(:NH)CET.HCl, m. 88° then PhCHEC(:NH)NH2.HCl, m.
236°, PhCHEPC(:NH)NHPP, m. 86°. Alao, PhCHEPC(:NH)NH2.E, b0.1
102° and PhCHPC(:NH)NHPP, m. 110.5°. PhCH(C8H17-n)CN gives
the amidines PhC(:SH17-n)C(:NH)NHPP, m. 150°. ThCH(C8H17-n)CN gives
the amidines PhC(:SH17-n)C(:NH)NHPP, m. 150°. ThCH(C8H17-n)CN gives
the amidines PhC(:SH17-n)C(:NH)NHPP, m. 150°. ThCH(C8H17-n)CN gives
the amidines PhC(:SH17-n)C(:NH)NHPP, m. 52°. The Physiol. activity of the
substituted amidines is being studied.

TO 78622-24-9 CAPLUS
CN Benzeneethanimidamide, α-methyl-, monohydrochloride (9CI) (CA INDEX
NAME)

Ph NH

● HC

RN 91429-53-7 CAPLUS CN Hydratropamidine, N-ethyl- (6CI, 7CI) (CA INDEX NAME)

Ph NH | || Me-CH-C-NHE

Page 39 10/568,760

ANSWER 59 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) 92579-12-9 CAPLUS Hydratropamidine, N-phenyl- (6CI, 7CI) (CA INDEX NAME)

HN Ph || | PhNH-C-CH-Me

ANSWER 60 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN (Continued) bi4 81-5°, np25 1.4922, and 22 g. α-phenylisobutyrohydroxamyl chloride (VIII), Me2PhCCC1:NOH, m. 73-4°. VIII and PhNB2 in EtoH give α-phenylisobutyrohydroxamanilide, m. 171-2°. VIII (5 g.) and 2.1 g. NaHcO3 in 100 cc. H2O, shaken 0.5 hr., give 2 g. 1-phenylisopropyl isocyanate (IX), b0.16 50-2°, np22 1.5038; with PhNNB2 it yields 1-phenyl-3 (-1-phenylisopropyl)urea, m. 193-4°. IX (0.6 g.), refluxed 3 hrs. with aq. NaHCO3, gives 0.4 g. 1,3-bis(1-phenylisopropyl)urea, (PhNe2CHN)2CO, m. 226-7°. Hydrospathon of 4 g. of IX over Raney Ni gives 1.6 g. 1. methyl-1,3-bis(1-phenylisopropyl)urea, PhMe2CHN9CCOMHC-PhMe2, m. 171-2°. MeCH(OH)CHNO2 (50 g.), C6H6, and AlCl3, refluxed 4 hrs., give 23% PhCH:NOH and 9 g. MeCHPh2; this can be explained by the initial formation of MeNO2 and MeCHO. MeNO2 and C6H6 give PhCH:NOH and PhN:CHPh. 858208-39-6P, Hydratropanilide, α-methyl-, oxime RI: PREP (Preparation) (preparation of) 858208-39-6 CAPLUS
Benzeneethanimidamide, N-hydroxy-α,α-dimethyl-N'-phenyl- (CA INDEX NAME)

Ph NH-OH L4 ANSWER 60 OF 60 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1949:38809 CAPLUS

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 43:38809 43:6993c-i,6994a-b

ORIGINAL REFERENCE NO.: 43:69930-1,6994a-b

TITLE: Aliphatic nitro compounds. XIX. Friedel-Crafts reactions with a- and P-nitro olefins

AUTHOR(S): Lambert, A.; Rose, J. D.; Weedon, B. C. L.

SOURCE: JOURNAL OF CHEMICAL SOCIETY (1949) 42-6

CODEN: JCSOA9; ISSN: 0368-1769

DOCUMENT TYPE: JOURNAL

LANGUAGE: Unavailable

AB cf. c.A. 42, 4917e. CH2:CMCCHENO2 (I) (10 g.), added (15 min.) to 16 g.

AlC13 in 50 cc. C6H6 at 30°, stirred 1 hr. at 30-40°, and poured onto concentrated HCl and ice, gives 9.55 g. 1-nitro-2-phenyl-2-methylpropane (II), 80.1 67-70°, nD23 1.5235; II results in 13-g. yield from PhMgBr (9.6 g. Mg) in 300 cc. ether on addition (1.5 hrs.) to

g. Me2C:CHNO2 in 300 cc. H2O at -5 to 0°, refluxing 0.5 hr. decomposing with 24 g. AcOH in 160 cc. H2O. Reduction of 4

MeOH over Raney Ni at room temperature and atmospheric pressure gives

MeOH over Raney Ni at room temperature and atmospheric pressure give 2.5 g. 2-phenyl-2-methylpropylamine, bl4 96-8° (picrate, yellow, m. 160°). I (10 g.) in 100 cc. PhMe, saturated with BF3, kept at room temperature overnight, and heated 1 hr. at 70-80°, gives 5 g. 1-nitro-2-p-tolyl-2-methylpropane (III), pale yellow, b0.5 85-90°, b13 145-50°, nD22 1.5258. III (0.4 g.), boiled 4 hrs. with 3 g. RMnO4 in 25 cc. H2O, gives 0.3 g. a,a-dimethylhometerephthatic acid (IV), m. 236-7°. Catalytic reduction (as above) of 3.6 g. III yields 2.5 g. 2-p-tolyl-2-methylpropylamine, b10 II1.5-15°, b22 134°, nD22 1.5231 (picrate, yellow, m. 211-13°). Me2CiNNO2 (20 g.) in 100 cc. PhMe, saturated with BF3 at 50°, gives 8.5 g. a-(p-tolyl)isobutyrohydroxamic acid (V), m. 157°, gives a deep red-violet color with FeCl3, and reduces AgNO3 in NH4OH. Distillation of

compound

the residue from the PhMe yields 6 g. III and a small quantity of a ound (CIIH130N ?), m. 132-4°. Catalytic reduction of 0.9 g. V in MeOH yields 0.5 g. α-(p-tolyl)lisobutyramide (VI), m. 143-4°. VI (0.2 g.) and 15 cc. 2 N HCl, refluxed 10 hrs., give 0.17 g. α-(p-tolyl)isobutyric acid (VII), m. 82°, VII results also on refluxing 0.5 g. V and 15 cc. 2 N HCl 0.5 hr. The structures of the VII reported by Wallach (Nachr. Ges. Wiss. Gottingen 2, 4(1899)] and by Rupe and Burgin (C.A. 5, 2841) are not clear. The Na derivative from 98 g. p-MeC6HGCHECKO (prepared with NaNNI2), treated dropwise with 213 g. MeI (1 hr.), gives 42% α-(p-tolyl)lisobutyronitrile, bl2 122-3°, b763 246°, np22 1.5106, d2222 0.9661; hydrolysis with KOH yields VII and VII. Oxidation of 4 g. VII in 40 cc. 5% Na2CO3 with 240 cc. 4% KNMO4 (1 hr.) gives 4 g. IVy further oxidation gives p-C6H4(CO2H)2. Details are given of the attempted preparation of VII by the method of R. and B. Me2C:CHNO2 and C6H6 do not yield a hydroxamic acid with BF3. Me2C:CHNO2 (60 g.), added (1 hr.) to 80 g. AlCl3 in 300 cc. C6H6 at 40°, the mixture stirred an addnl. hr., decomposed with HCl and ice, and acted with a time 2 for the stirred an addnl. hr., decomposed with HCl and ice, and acted with a crise 2 for a flatiplicy solutive advance of the stirre 2 for a flatiplicy solutive advance and well acted with SCH6 crises 2 for a flatiplicy solutive advance we decoded to the collection of the stirred an addnl. hr., decomposed with HCl and ice, and acted with

C6H6, gives 25 g. α, β -dichloroisobutyraldoxime, Me2CClCCl:NOH,